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(54) Title: IDENTIFICATION OF SORTASE GENE

(57) Abstract

The present invention substantially purified sortase-transamidase enzyme from Gram-positive bacteria, such as Staphylococcus aureus. The enzyme having a molecular weight of about 41,000 daltons and catalyzing a reaction that covalently cross-links the carboxyl terminus of a protein having a sorting signal to the peptidoglycan of a Gram-positive bacterium, the sorting signal having: (1) a motif of LPX₃X₄G therein; (2) a substantially hydrophobic domain of at least 31 amino acids carboxyl to the motif; and (3) a charged tail region with at least two positively charged residues carboxyl to the substantially hydrophobic domain, at least one of the two positively charged residues being arginine, the two positively charged residues being located at residues 31-33 from the motif, wherein X₃ is any of the twenty naturally-occurring L-amino acids and X4 is selected from the group consisting of alanine, serine, and threonine, and wherein sorting occurs by cleavage between the fourth and fifth residues of the

LPX₃X₄G motif. Variants of the enzyme, methods for cloning the gene encoding the enzyme and expressing the cloned gene, and methods of use of the enzyme, including for screening for antibiotics and for display of proteins or peptides on the surfaces of Gram-positive bacteria, are also disclosed.

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IDENTIFICATION OF SORTASE GENE

GOVERNMENT RIGHTS

This invention was supported by grants from the United States government, namely grants from the National Institutes of Health, NIH-NIAID Grant Nos. AI 33985 and 38897. Accordingly, the government may have certain rights in this invention.

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BACKGROUND OF THE INVENTION

This invention is directed to an enzyme from Gram-positive bacteria, designated sortase-transamidase, nucleic acid segments encoding the enzyme, and methods of use of the enzyme.

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Human infections caused by Gram-positive bacteria present a medical challenge due to the dramatic increase in multiple antibiotic resistance strains in recent years. Gram-positive bacteria that can cause serious or fatal infections in humans include Staphylococcus, Streptococcus, Enterococcus, Pneumococcus, Bacillus, Actinomyces, Mycobacterium, and Listeria, as well as others. Infections caused by these pathogens are particularly severe and difficult to treat in immunologically compromised patients. These include patients suffering from infection with the Human Immunodeficiency Virus (HIV), the virus that causes AIDS, as well as patients given immune-suppressive agents for treatment of cancer or autoimmune diseases. In particular, infections caused by various Mycobacterium species, including M. tuberculosis, M. bovis, M. avium, and M. intracellulare, are frequently the cause of disease in patients with AIDS.

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Therefore, it is apparent that new target sites for bacterial chemotherapy are needed if such pathogenic organisms are to be controlled.

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A unique characteristic of these pathogens and many Gram-positive bacteria is their surface display of proteins anchored to the cell wall. In fact, many of these molecules are known to be involved in essential cellular functions, including pathogenesis in a susceptible host. Thus, a possible disruption in this anchoring process may prove to be an effective treatment against these disease—causing elements.

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The anchoring of surface molecules to the cell wall in Gram-positive bacteria has been demonstrated to involve a conserved pathway, culminating in recognition of a conserved cleavage/anchoring site by some previously uncharacterized cellular machinery. Molecules whose ultimate location is the cell

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wall must invariably be translocated across the single cellular membrane of these organisms. This is mediated for all cell wall anchored proteins by the well studied secretory pathway, involving cleavage of an amino-terminal signal peptide by a type I signal peptidase. Upon translocation of the molecule out of the cytoplasm, a mechanism must be present that extracellularly recognizes this protein as a substrate for anchoring. This process has been previously shown to involve the carboxyl-terminally located cell wall sorting signal, consisting of a highly conserved motif such as LPXTG (SEQ ID NO:1), in which X can represent any of the twenty naturally occurring L-amino acids, followed by a series of hydrophobic residues and ultimately a sequence of positively-charged residues. Thus, once amino-terminally modified and successfully secreted, a polypeptide with this carboxyl-terminal sequence can present itself as a substrate to be processed by the anchoring machinery. At this time, cleavage of the sorting signal after the threonine residue is coupled with covalent linkage of the remainder of the polypeptide to the free amino group of the pentaglycine crossbridge in the cell wall.

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It is this transpeptidation reaction that anchors mature surface proteins so that the peptidoglycan layer, from which point the molecules can serve their biological functions. Therefore, there is a need to isolate and purify the enzyme that catalyzes this reaction. There is also a need to identify the gene encoding such an enzyme in order that the enzyme can be produced by genetic engineering techniques.

Additionally, there is also a need to develop new methods for displaying proteins or peptides on the surfaces of bacteria. For many purposes, it is desirable to display proteins or peptides on the surfaces of bacteria so that the proteins or peptides are accessible to the surrounding solution, and can, for example, be bound by a ligand that is bound specifically by the protein or peptide. In particular, the display of proteins on the surface of bacteria is desirable for the preparation of vaccines, the linkage of molecules such as antibiotic molecules or diagnostic reagents to cells, for screening reagents such as monoclonal antibodies, and for the selection of cloned proteins by displaying the cloned proteins, then observing their reaction with specific reagents such as antibodies. One way of doing this has been with phage display (G.P. Smith, "Filamentous Fusion Phage: Novel Expression Vectors that Display Cloned Antigens on the Virion Surface," Science 228:1315-1316 (1985)). However, phage display is limited in its practicality, because it requires that the protein being displayed to be inserted into a coat protein of filamentous phage and retain its activity while not distorting the conformation of the coat protein, allowing functional virions to be formed. In general, this technique is therefore limited only to small peptide and proteins.

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Therefore, there is a need for a more general method of peptide and protein display.

SUMMARY

The present invention is directed to sortase-transamidase enzymes from Gram-positive bacteria, particularly Staphylococcus aureus, and methods for their use, particularly in the areas of drug screening and peptide and protein display.

One aspect of the present invention is a substantially purified sortasetransamidase enzyme from a Gram-positive bacterium, the enzyme catalyzing a reaction that covalently cross-links the carboxyl terminus of a protein having a sorting signal to the peptidoglycan of a Gram-positive bacterium, the sorting signal having a motif of LPX₃X₄G therein, wherein sorting occurs by cleavage between the fourth and fifth residues of the LPX₃X₄G motif. Typically, the Gram-positive bacterium is a species selected from the group consisting of Staphylococcus aureus, S. sobrinus. Enterococcus faecalis, Streptococcus pyogenes, and Listeria monocytogenes.

Preferably, the Gram-positive bacterium is S. aureus. The enzyme may be a heterooligomer.

Preferably, the enzyme has at least one subunit with a molecular weight of about 41,000 daltons and the sorting signal further includes: (2) a substantially hydrophobic domain of at least 31 amino acids carboxyl to the motif; and (3) a charged tail region with at least two positively charged residues carboxyl to the substantially hydrophobic domain, at least one of the two positively charged residues being arginine, the two positively charged residues being located at residues 31-33 from the motif, wherein X3 is any of the twenty naturally-occurring L-amino acids and X₄ is selected from the group consisting of alanine, serine, and threonine.

Preferably, the enzyme includes therein a subunit whose amino acid sequence is selected from the group consisting of: (1) D-P-K-L-K-E-I-Y-O-I-V-E-S-K-G-Y-G-K-E-F-G-H-S-L-G-H-G-I-G-L-E-I-H-E-G-P-M-L-A-D-D-I-L-I-T-E-N-G-C-Q-V-F-T-K-C-T-K-D-L-I-V-L-T (SEQ ID NO: 2); (2) M-V-K-V-T-D-Y-S-N-S-K-L-G-K-E-I-A-P-E-V-L-S-V-I-A-S-I-A-T-S-E-V-E-G-I-T-G-H-F-A-E-L-K-E-T-N-L-E-K-V-S-R-K-N-L-S-R-D-L-K-I-E-S-K-E-G-I-Y-I-D-V-Y-C-A-L-K-H-G-V-N-I-S-K-T-A-N-K-I-Q-T-S-I-F-N-S-I-S-N-M-T-A-I-E-P-K-Q-I-N-I-H-I-T-O-I-V-I-E-K (SEQ ID NO: 31) and (3) sequences incorporating one or more conservative amino acid substitutions in SEQ ID NO:2 or SEQ ID NO: 31, wherein

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the conservative amino acid substitutions are any of the following: (1) any of isoleucine, leucine, and valine for any other of these amino acids; (2) aspartic acid for glutamic acid and vice versa; (3) glutamine for asparagine and vice versa; and (4) serine for threonine and vice versa.

Another aspect of the present invention is a nucleic acid sequence encoding this enzyme. In one alternative, the nucleic acid sequence includes therein a sequence selected from the group consisting of: (1)

GATCCTAAACTGAAAGAAATATATCAAATAGTACTTGAATCTCAAATGAA
AGCAATTAATGAGATTAGACCTGGCATGACTGGTGCAGAAGCTGATGCCA
TTTCAAGAAACTATTTAGAGTCAAAAGGGTATGGAAAAGAATTTGGACAT
TCACTAGGACATGGTATTGGTTTAGAAATCCATGAAGGGCCAATGCTGGC
TCGTACGATACAAGATAAACTTCAAGTTAACAACTGTGTTACAGTAGAAC
CTGGTGTTTATATAGAAGGTTTGGGCGGTATAAGAATAGAAGAAGACCTT
ATAGTTTAACATAA (SEQ ID NO: 28); (2)
ATGGTCAAAGTAACTGATTATTCAAATTCAAAATTAGGTAAAAGTAGAAAT
AGCGCCAGAAGTGCTATCTGTTATTGCAAGTATAGCTACTTCGGAAGTCG
AAGGCATCACTGGCCATTTTTTTTTAAAAAAAAATTTAGAAAAAA

AAGCATCACTGGCCATTTTGCTGAATTAAAAGAAACAAATTTAGAAAAA GTTAGTCGTAAAATTTAAGCCGTGATTTAAAAAATTTAGAAAAAA GTTAGTCGTAAAAAATTTAAGCCGTGATTTAAAAAATCGAGAGTAAAGAAGA TGGCATATATATAGATGTATATTGTGCATTAAAAACATGGTAATATTTCAAA AACTGCAAACAAAATTCAAACGTCAATTTTTAATTCAATTTCTAATATGAC AGCGATAGAACCTAAGCAAATTAATATTCACATTACACAAATCGTTATTG AAAAGTAA (SEQ ID NO: 30); and (3) a sequence complementary to SEQ ID NO: 28 or SEQ ID NO: 30. In another alternative, the nucleic acid sequence can include a sequence hybridizing with SEQ ID NO: 28, SEQ ID NO: 30 or a sequence complementary to SEQ ID NO: 28 or SEQ ID NO: 30 with no greater than about a 15% mismatch under stringent conditions. Preferably, the degree of mismatch is less than about 5%; more preferably, the degree of mismatch is less than about 2%.

Yet another aspect of the present invention is a vector comprising the nucleic acid sequence of the present invention operatively linked to at least one control sequence that controls the expression or regulation of the nucleic acid sequence.

Yet another aspect of the present invention is a host cell transfected with a vector of the present invention.

Another aspect of the present invention is a method for producing a substantially purified sortase-transamidase enzyme. The method comprises the steps of:

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- (1) culturing a host cell according to the present invention under conditions in which the host cell expresses the encoded sortase-transamidase enzyme; and
- (2) purifying the expressed enzyme to produce substantially purified sortase-transamidase enzyme.

Another aspect of the present invention is a method for screening a compound for anti-sortase-transamidase activity. This method is important in providing a way to screen for antibiotics that disrupt the sorting reaction and are likely to be effective in treating infections caused by Gram-positive bacteria.

In one alternative, the screening method comprises the steps of:

- (1) providing a substantially purified sortase—transamidase enzyme according to the present invention;
- (2) performing an assay for sortase-transamidase in the presence and in the absence of the compound; and
- (3) comparing the activity of the sortase—transamidase enzyme in the presence and in the absence of the compound to screen the compound for sortase—transamidase activity.

In another alternative, the screening method comprises the steps of:

- (1) providing an active fraction of sortase-transamidase enzyme from a Gram-positive bacterium;
 - (2) performing an assay for sortase-transamidase in the presence and in the absence of the compound; and
 - (3) comparing the activity of the sortase-transamidase enzyme in the presence and in the absence of the compound to screen the compound for sortase-transamidase activity.

The active fraction of sortase-transamidase activity can be a particulate fraction from *Staphylococcus aureus*.

The assay for sortase—transamidase enzyme can be performed by monitoring the capture of a soluble peptide that is a substrate for the enzyme by its interaction with an affinity resin. In one alternative, the soluble peptide includes a sequence of at least six histidine residues and the affinity resin contains nickel. In another alternative, the soluble peptide includes the active site of glutathione S—transferase and the affinity resin contains glutathione. In yet another alternative, the soluble peptide includes the active site of streptavidin and the affinity resin contains biotin. In still another alternative, the soluble peptide includes the active site of maltose binding protein and the affinity resin contains amylose.

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Still another aspect of the present invention is an antibody specifically binding the sortase—transamidase enzyme of the present invention.

Yet another aspect of the present invention is a protein molecule comprising a substantially purified sortase—transamidase enzyme according to the present invention extended at its carboxyl—terminus with a sufficient number of histidine residues to allow specific binding of the protein molecule to a nickel—sepharose column through the histidine residues added at the carboxyl—terminus.

Still another aspect of the present invention is a method for displaying a polypeptide on the surface of a Gram-positive bacterium comprising the steps of:

- (1) expressing a polypeptide having a sorting signal at its carboxy-terminal end, the sorting signal having: (a) a motif of LPX₃X₄G therein; (b) a substantially hydrophobic domain of at least 31 amino acids carboxyl to the motif; and (c) a charged tail region with at least two positively charged residues carboxyl to the substantially hydrophobic domain, at least one of the two positively charged residues being arginine, the two positively charged residues being located at residues 31-33 from the motif, wherein X₃ is any of the twenty naturally-occurring L-amino acids and X₄ is selected from the group consisting of alanine, serine, and threonine;
- (2) forming a reaction mixture including: (i) the expressed polypeptide; (ii) a substantially purified sortase-transamidase according to the present invention; and (iii) a Gram-positive bacterium having a peptidoglycan to which the sortase-transamidase can link the polypeptide; and
- (3) allowing the sortase-transamidase to catalyze a reaction that cleaves the polypeptide within the LPX₃X₄ motif of the sorting signal and covalently cross-links the amino-terminal portion of the cleaved polypeptide to the peptidoglycan to display the polypeptide on the surface of the Gram-positive bacterium.

Another display method according to the present invention comprises:

- (1) cloning a nucleic acid segment encoding a chimeric protein into a Gram-positive bacterium to generate a cloned chimeric protein including therein a carboxyl-terminal sorting signal as described above;
- (2) growing the bacterium into which the nucleic acid segment has been cloned to express the cloned chimeric protein to generate a chimeric protein including therein a carboxyl-terminal sorting signal; and
- (3) binding the polypeptide covalently to the cell wall by the enzymatic action of a sortase-transamidase expressed by the Gram-positive bacterium involving cleavage of the chimeric protein within the LPX₃X₄G motif so that the polypeptide is

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displayed on the surface of the Gram-positive bacterium in such a way that the polypeptide is accessible to a ligand.

Another aspect of the present invention is a polypeptide displayed on the surface of a Gram-positive bacterium by covalent linkage of an amino-acid sequence of LPX₃X₄ derived from cleavage of an LPX₃X₄G motif, wherein X₃ is any of the twenty naturally-occurring L-amino acids and X₄ is selected from the group consisting of alanine, serine, and threonine, the polypeptide being displayed on the surface of the Gram-positive bacterium in such a way that the polypeptide is accessible to a ligand.

Another aspect of the present invention is a covalent complex comprising:

- (1) the displayed polypeptide; and
- (2) an antigen or hapten covalently cross-linked to the polypeptide.

Yet another aspect of the present invention is a method for vaccination of an animal comprising the step of immunizing the animal with the displayed polypeptide to generate an immune response against the displayed polypeptide, or, alternatively, with the covalent complex to generate an immune response against the antigen or the hapten.

Still another aspect of the present invention is a method for screening for expression of a cloned polypeptide comprising the steps of:

- (1) expressing a cloned polypeptide as a chimeric protein having a sorting signal at its carboxy-terminal end as described above;
- (2) forming a reaction mixture including: (i) the expressed chimeric protein; (ii) a substantially purified sortase—transamidase enzyme according to the present invention; and (iii) a Gram—positive bacterium having a peptidoglycan to which the sortase—transamidase can link the polypeptide through the sorting signal;
- (3) binding the chimeric protein covalently to the cell wall by the enzymatic action of a sortase-transamidase expressed by the Gram-positive bacterium involving cleavage of the chimeric protein within the LPX₃X₄G motif so that the polypeptide is displayed on the surface of the Gram-positive bacterium in such a way that the polypeptide is accessible to a ligand; and
- (4) reacting the displayed polypeptide with a labeled specific binding partner to screen the chimeric protein for reactivity with the labeled specific binding partner.
- Still another aspect of the present invention is a method for the diagnosis or treatment of a bacterial infection caused by a Gram-positive bacterium comprising the steps of:

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(1) conjugating an antibiotic or a detection reagent to a protein including therein a carboxyl-terminal sorting signal as described above to produce a conjugate; and

(2) introducing the conjugate to an organism infected with a Grampositive bacterium in order to cause the conjugate to be sorted and covalently crosslinked to the cell walls of the bacterium in order to treat or diagnose the infection.

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If an antibiotic is used, typically it is a penicillin, ampicillin, vancomycin, gentamicin, streptomycin, a cephalosporin, amikacin, kanamycin, neomycin, paromomycin, tobramycin, ciprofloxacin, clindamycin, rifampin, chloramphenicol, norfloxacin, or a derivative of these antibiotics.

Similarly, another aspect of the present invention is a conjugate comprising an antibiotic or a detection reagent covalently conjugated to a protein including therein a carboxyl—terminal sorting signal as described above to produce a conjugate. In still another aspect of the present invention, a composition comprises the conjugate with a pharmaceutically acceptable carrier.

Another aspect of the present invention is a substantially purified protein having at least about 50% match with best alignment with the amino acid sequences of at least one of the putative *Bacillus* peptidase (SEQ ID NO: 3), the aminopeptidase P of *Lactococcus lactis* (SEQ ID NO: 4), or the proline dipeptidase of *Lactobacillus delbrueckii lactis* (SEQ ID NO: 5) and having sortase—transamidase activity. Preferably, the match is at least about 60% in best alignment; more preferably, the match is at least about 70% in best alignment.

Another aspect of the present invention is a substantially purified protein having sortase—transamidase activity and a hydrophobicity profile of at least one subunit of the protein that, determined as the mean absolute value of the hydrophobicity difference per residue, differs from the hydrophobicity profile of a putative *Bacillus* peptidase (SEQ ID NO: 3) by no more than about 2 units on the hydrophobicity scale. Preferably, the difference is not more than about 1 unit; most preferably, it is not more than about 0.5 units.

Another aspect of the present invention is a substantially purified protein having sortase—transamidase activity and a hydrophobicity profile of at least one subunit of the protein that, determined as the mean absolute value of the hydrophobicity difference per residue, differs from the hydrophobicity profile of the sequence of SEQ ID NO: 31 by no more than about 2 units on the hydrophobicity scale.

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BRIEF DESCRIPTION OF THE DRAWINGS

These and other features, aspects, and advantages of the present invention will become better understood with reference to the following description and accompanying drawings where:

Figure 1 is a diagram showing the substantial homology of the amino acid sequence of the sortase-transamidase enzyme of *Staphylococcus aureus* to an open reading frame in the genome of *Streptococcus pyogenes* (SEQ ID NO: 2 & 34);

Figure 2 is a diagram showing a greater homology of the amino acid sequence of the sortase—transamidase enzyme of *Staphylococcus aureus* to the carboxyl—terminal portion of the open reading frame in the genome of *Streptococcus pyogenes* (SEQ ID NO: 2 & 34);

Figure 3 is the DNA sequence of the S. pyogenes open reading frame (SEQ ID NO: 33 & 34);

Figure 4 (SEQ ID NO: 34) is the entire amino acid sequence of the protein translated from the entire S. pyogenes open reading frame;

Figure 5 (SEQ ID NO: 3) is the amino acid sequence of a putative *Bacillus* peptidase in the GCVT-SPOIIIAA intergenic region;

Figure 6 is the hydrophobicity profile of the protein whose amino acid sequence is shown in Figure 5 (SEQ ID NO: 3);

Figure 7 (SEQ ID NO: 4) is the amino acid sequence of the aminopeptidase P of *Lactococcus lactis*;

Figure 8 (SEQ ID NO: 5) is the amino acid sequence of the proline dipeptidase of *Lactobacillus delbrueckii lactis*;

Figure 9 is a diagram of the activity of the sortase—transamidase enzyme of the present invention;

Figure 10 (SEQ ID NOS: 28 & 29) is a partial DNA sequence of the gene for one of the subunits of the sortase-transamidase enzyme of S. aureus;

Figure 11 (SEQ ID NO: 2) is the partial carboxyl-terminal amino acid sequence translated from the DNA sequence of Figure 10 (SEQ ID NOS: 28 & 29);

Figure 12 (SEQ ID NOS: 30 & 31) is a partial DNA sequence of the gene for a second of the subunits of the sortase—transamidase enzyme of S. aureus; and

Figure 13 is the hydrophobicity profile of the protein translated from the DNA sequence of Figure 12 (SEQ ID NOS: 30 & 31).

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DEFINITIONS

As used herein, the terms defined below have the following meanings unless otherwise indicated:

"Nucleic Acid Sequence": the term "nucleic acid sequence" includes both DNA and RNA unless otherwise specified, and, unless otherwise specified, includes both double—stranded and single—stranded nucleic acids. Also included are hybrids such as DNA—RNA hybrids. In particular, a reference to DNA includes RNA that has either the equivalent base sequence except for the substitution of uracil and RNA for thymine in DNA, or has a complementary base sequence except for the substitution of uracil for thymine, complementarity being determined according to the Watson—Crick base pairing rules. Reference to nucleic acid sequences can also include modified bases as long as the modifications do not significantly interfere either with binding of a ligand such as a protein by the nucleic acid or with Watson—Crick base pairing.

"Antibody": as used herein the term "antibody" includes both intact antibody molecules of the appropriate specificity, and antibody fragments (including Fab, F(ab'), Fv, and F(ab')₂), as well as chemically modified intact antibody molecules and antibody fragments, including hybrid antibodies assembled by in vitro reassociation of subunits. Also included are single—chain antibody molecules generally denoted by the term sFv and humanized antibodies in which some or all of the originally non—human constant regions are replaced with constant regions originally derived from human antibody sequences. Both polyclonal and monoclonal antibodies are included unless otherwise specified. Additionally included are modified antibodies or antibodies conjugated to labels or other molecules that do not block or alter the binding capacity of the antibody.

DESCRIPTION

A substantially purified sortase-transamidase enzyme from Gram-positive bacteria, particularly *Staphylococcus aureus*.

The properties of this enzyme make it a logical target for antibiotic action. This enzyme also catalyzes covalent crosslinkage of proteins to the peptidoglycan of Gram-positive bacteria.

I. THE SORTASE-TRANSAMIDASE ENZYME

One aspect of the invention is a substantially purified sortase transamidase enzyme from a Gram—positive bacterium. As used herein, the term "substantially purified" means having a specific activity of at least tenfold greater than the sortase-transamidase activity present in a crude extract, lysate, or other state from which proteins have not been removed and also in substantial isolation from proteins found in association with sortase-transamidase in the cell.

One subunit of the enzyme has a molecular weight of about 41,000 daltons. The enzyme catalyzes a reaction that covalently crosslinks the carboxylterminus of a protein having a sorting signal to the peptidoglycan of the Grampositive bacterium. The sorting signal has: (1) a motif of LPX₃X₄G therein; (2) a substantially hydrophobic domain of at least 31 amino acids carboxyl to the motif; and (3) a charged tail region with at least two positively charged residues carboxyl to the substantially hydrophobic domain, at least one of the two positively charged residues being arginine, the two positively charged residues being located at residues 31-33 from the motif. In this sorting signal, X₃ can be any of the twenty naturally-occurring L-amino acids. X₄ can be alanine, serine, or threonine. Preferably, X₄ is threonine.

The sortase-transamidase is believed to occur in all Gram-positive bacteria. In particular, the enzyme exists in Mycobacterium, Nocardia, Actinomyces. Staphylococcus, Streptococcus, Listeria, Enterococcus, and Pneumococcus. Specifically, the enzyme exists in the following species: Staphylococcus aureus, S. sobrinus, Enterococcus faecalis, Streptococcus pyogenes, and Listeria monocytogenes.

Preferably the enzyme is isolated from Staphylococcus aureus.

A. Amino Acid Sequence

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The sortase-transamidase of the present invention includes therein an amino acid sequence, in one subunit of the enzyme, of D-P-K-L-K-E-I-Y-O-I-V-L-E-S-Q-M-K-A-I-N-E-I-R-P-G-M-T-G-A-E-A-D-A-I-S-R-N-Y-L-E-S-K-G-Y-G-K-E-F-G-H-S-L-G-H-G-I-G-L-E-I-H-E-G-P-M-L-A-R-P-I-Q-D-K-L-Q-V-N-N-C-V-T-V-E-P-G-V-Y-I-E-G-L-G-G-I-R-I-E-D-D-I-L-I-T-E-N-G-C-Q-V-F-T-K-C-T-K-D-L-I-V-L-T (SEO ID NO:2). This sequence is at the carboxyl-terminal end of the subunit of the enzyme.

The sortase-transamidase of the present invention also includes therein an amino acid sequence, in a second subunit of the enzyme, of M-V-K-V-T-D-Y-S-N-S-K-L-G-K-E-I-A-P-E-V-L-S-V-I-A-S-I-A-T-S-E-V-E-G-I-T-G-H-F-A-E-L-K-E-T-N-L-E-K-V-S-R-K-N-L-S-R-D-L-K-I-E-S-K-E-G-I-Y-I-D-V-Y-C-A-L-K-H-G-V-N-I-S-K-T-A-N-K-I-O-T-S-I-F-N-S-I-S-N-M-T-A-I-E-P-K-Q-I-N-I-H-I-T-Q-I-V-I-E-K (SEO ID NO: 31).

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Also within the scope of the present invention are substantially purified protein molecules that are mutants of the sequence of SEQ ID NO: 31 that preserve the sortase—transamidase activity. In particular, the conservative amino acid substitutions can be any of the following: (1) any of isoleucine, leucine, and valine for any other of these amino acids; (2) aspartic acid for glutamic acid and vice versa; (3) glutamine for asparagine and vice versa; and (4) serine for threonine and vice versa.

Other substitutions can also be considered conservative, depending upon the environment of the particular amino acid. For example, glycine (G) and alanine (A) can frequently be interchangeable, as can be alanine and valine (V). Methionine (M), which is relatively hydrophobic, can frequently be interchanged with leucine and isoleucine, and sometimes with valine. Lysine (K) and arginine (R) are frequently interchangeable in locations in which the significant feature of the amino acid residue is its charge and the different pK's of these two amino acid residues or their different sizes are not significant. Still other changes can be considered "conservative" in particular environments. For example, if an amino acid on the surface of a protein is not involved in a hydrogen bond or salt bridge interaction with another molecule, such as another protein subunit or a ligand bound by the protein, negatively charged amino acids such as glutamic acid and aspartic acid can be substituted for by positively charged amino acids such as lysine or arginine and vice versa. Histidine (H), which is more weakly basic than arginine or lysine, and is partially charged at neutral pH, can sometimes be substituted for these more basic amino acids. Additionally, the amides glutamine (O) and asparagine (N) can sometimes be substituted for their carboxylic acid homologues, glutamic acid and aspartic acid.

The sortase-transamidase from Staphylococcus aureus has substantial homology the amino acid sequence of the first subunit, that of SEQ ID NO: 2, with an open reading frame in the genome of Streptococcus pyogenes, particularly in the amino-terminal region. There is about a 22% match with best alignment over the entire sequenced region of the S. pyogenes open reading frame, and about a 47% match with best alignment over the carboxyl-terminal region of the S. pyogenes open reading frame. These matches are shown in Figures 1-2. The DNA sequence of the entire S. pyogenes open reading frame is shown in Figure 3 (SEQ ID NO: 33 & 34). The protein translated from the entire S. pyogenes open reading frame has a molecular weight of about 40,851.43 daltons; its sequence is shown in Figure 4 (SEQ ID NO: 34). Therefore, another aspect of the present invention is a substantially purified protein molecule that has at least one subunit of about 40,000 to about 41,000 daltons

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in molecular weight, the subunit having at least a 20% match with best alignment with the S. pyogenes open reading frame of Figure 2 and that has sortase—transamidase activity. Preferably, the subunit has at least a 30% match with best alignment; more preferably, at least 50% match with best alignment.

As shown below in the Example, the first 364 bases of a nucleic acid segment that complements a temperature—sensitive mutation in the *S. aureus* sortase—transamidase, designated the SM-317 complementing gene insert, has been identified as encoding a protein sequence that is a homologue of a putative *Bacillus* peptidase in the GCVT-SPOIIIAA intergenic region (GenBank Accession No. 1731048; Y. Kobayashi et al.). The sequence of this putative peptidase is shown in Figure 5 (SEQ ID NO:3) and its hydrophobicity profile is shown in Figure 6. The hydrophobicity is calculated according to the method of J. Kyte & R.F. Doolittle, "A Simple Method for Displaying the Hydropathic Character of a Protein," J. Mol. Biol. 157: 105-132

(1982). As used herein, the term "hydrophobicity" is the hydrophobicity as calculated

in Kite & Doolittle, supra.

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To a lesser degree of homology, the protein sequence encoded by this complementing gene insert is homologous to aminopeptidase P of Lactococcus lactis (GenBank Accession No. 1915907; J. Matos). The amino acid sequence of this aminopeptidase is shown in Figure 7 (SEQ ID NO: 4). To a still lesser degree of homology, the protein sequence encoded by this complementing gene insert is homologous to the proline dipeptidase of Lactobacillus delbrueckii lactis (GenBank Accession No. 1172066; K. Stucky et al., "Cloning and DNA Sequence Analysis of pepQ, a Prolidase Gene from Lactobacillus delbrueckii subsp. lactis and Partial Characterization of Its Product," Mol. Gen. Genet. 247: 494–500 (1995)). The amino acid sequence of this proline dipeptidase is shown in Figure 8 (SEO ID NO:5).

Because of the relatedness of these proteins, another aspect of the present invention is a substantially purified protein having at least one subunit with at least about 50% match with best alignment with the amino acid sequences of at least one of the putative *Bacillus* peptidase (SEQ ID NO: 3), the aminopeptidase P of *Lactococcus lactis* (SEQ ID NO: 4), or the proline dipeptidase of *Lactobacillus delbrueckii lactis* (SEQ ID NO: 5) and having sortase—transamidase activity. Preferably, the at least one subunit of the protein has at least about 60% match with best alignment with at least one of these sequence; more preferably, the at least one subunit of the protein has at least about 70% match with best alignment with at least one of these sequences.

Because the hydrophobicity of a protein is a sensitive measure of protein structure, another aspect of the invention is a substantially purified protein

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having sortase—transamidase activity and a hydrophobicity profile for at least one subunit of the protein that, determined as the mean absolute value of the hydrophobicity difference per residue, differs from the hydrophobicity profile of the putative *Bacillus* peptidase by no more than about 2 units on the hydrophobicity scale of Kyte & Doolittle, <u>supra</u>. Preferably, the difference is no greater than about 1 unit; more preferably, the difference is no greater than about 0.5 units.

The sortase-transamidase is a cysteine protease.

B. Activity of the Sortase-Transamidase

The activity of the sortase–transamidase enzyme of the present invention is shown, in general, in Figure 9. The enzyme first cleaves a polypeptide having a sorting signal within the LPX $_3$ X $_4$ G motif. Cleavage occurs after residue X $_4$, normally a threonine; as indicated above, this residue can also be a serine or alanine residue. This residue forms a covalent intermediate with the sortase. The next step is the transamidation reaction that transfers the cleaved carboxyl terminus of the protein to be sorted to the -NH $_2$ of the pentaglycine crossbridge within the peptidoglycan precursor. The peptidoglycan precursor is then incorporated into the cell wall by a transglycosylase reaction with the release of undecaprenyl phosphate. The mature anchored polypeptide chains are thus linked to the pentaglycine cross bridge in the cell wall which is tethered to the ε -amino side chain of an unsubstituted cell wall tetrapeptide. A carboxypeptidase may cleave a D-Ala-D-Ala bond of the pentapeptide structure to yield the final branched anchor peptide in the staphylococcal cell wall.

The sorting signal has: (1) a motif of LPX₃X₄G therein; (2) a substantially hydrophobic domain of at least 31 amino acids carboxyl to the motif; and (3) a charged tail region.

In the motif, X₃ can be any of the 20 naturally-occurring L-amino acids. X₄ can be any of threonine, serine, or alanine. Preferably, X₄ is threonine (O. Schneewind et al., "Cell Wall Sorting Signals in Surface Proteins of Gram-Positive Bacteria," <u>EMBO J.</u> 12:4803-4811 (1993)).

Preferably, the substantially hydrophobic domain carboxyl to the motif includes no more than about 7 charged residues or residues with polar side chains. For the purposes of this specification, these residues include the following: aspartic acid, glutamic acid, lysine, and arginine as charged residues, and serine, threonine, glutamine, and asparagine as polar but uncharged residues. Preferably, the sequence includes no more than three charged residues.

Representative sequences suitable for sorting signals for use with the sortase—transamidase of the present invention include, but are not limited to the following: E-E-N-P-F-I-G-T-T-V-F-G-G-L-S-L-A-L-G-A-A-L-L-A-G (SEQ ID NO: 6), the hydrophobic domain of the staphylococcal proteinase (SPA) sorting signal from *Staphylococcus aureus*; (2) G-E-E-S-T-N-K-G-M-L-F-G-G-L-F-S-I-L-G-L-A-L-L (SEQ ID NO:7), the SNBP signal of *S. aureus*; (3) D-S-S-N-A-Y-L-P-L-L-G-L-V-S-L-T-A-G-F-S-L-L-G-L (SEQ ID NO: 8), the SPAA signal of *S. sobrinus*, (4) E-K-Q-N-V-L-L-T-V-V-G-S-L-A-A-M-L-G-L-A-G-L-G-F (SEQ ID NO:9), the PRGB signal of *Enterococcus faecalis*, (5) S-I-G-T-Y-L-F-K-I-G-S-A-A-M-I-G-A-I-G-I-Y-I-V (SEQ ID NO:10), the TEE signal of *Streptococcus pyogenes*, and (6) D-S-D-N-A-L-Y-L-L-G-L-L-A-V-G-T-A-M-A-L-T (SEQ ID NO:11), the INLA signal of *Listeria monocytogenes*. Other hydrophobic domains can be used as part of the sorting signal.

The third portion of the sorting signal is a charged tail region with at least two positively charged residues carboxyl to the substantially hydrophobic 15 domain. At least one of the two positively charged residues is arginine. The charged tail can also contain other charged amino acids, such as lysine. Preferably, the charged tail region includes two or more arginine residues. The two positively charged residues are located at residues 31-33 from the motif. Preferably, the two arginine residues are either in succession or are separated by no more than one 20 intervening amino acid. Preferably, the charged tail is at least five amino acids long, although four is possible. Among the charged tails that can be used are the following: (1) R-R-R-E-L (SEQ ID NO:12), from the SPA signal of S. aureus; (2) R-R-N-K-K-N-H-K-A (SEQ ID NO:13), from the SNBP signal of S. aureus; (3) R-R-K-O-D (SEQ ID NO:14), from the SPAA signal of S. sobrinus; (4) K-R-R-K-E-T-K (SEQ ID NO:15), from the PRGB signal of E. faecalis; (5) K-R-R-K-A (SEQ ID NO:16), from the TEE signal of S. pyogenes; (6), K-R-R-H-V-A-K-H (SEO ID NO:17), from the FIM sorting signal of Actinomyces viscosus, and (7) K-R-R-K-S (SEQ ID NO:18), from the BAC sorting signal of Streptococcus aglactiae; (8) K-R-K-E-E-N (SEQ ID NO:19), from the EMM signal of Streptococcus pyogenes. 30

Also usable as the charged tail portion of the sorting signal are the following sequences produced by mutagenesis from the SPA signal of *S. aureus*. These include R-R-R-E-S (SEQ ID NO: 20), R-R-R-S-L (SEQ ID NO: 21), R-R-S-E-L (SEQ ID NO: 22), R-S-R-E-L (SEQ ID NO: 23) and S-R-R-E-L (SEQ ID NO: 24). Other charged tails that are usable as part of the sorting signal can be derived from a polyserine tail, itself inactive, by replacement of one or more of the serine residues with the basic amino acid arginine. These include R-R-S-S-S (SEO

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ID NO: 25), R-S-R-S-S (SEQ ID NO:26), and S-R-R-S-S (SEQ ID NO:27). Other sorting signals can also be used.

II. THE GENE ENCODING THE SORTASE-TRANSAMIDASE ENZYME

A. Isolation of the Sortase-Transamidase Enzyme Gene

The gene for the sortase-transamidase enzyme in Staphylococcus aureus has been isolated. The isolation process is described in detail in the Example below; in general, this process comprises: (1) the generation of temperature—sensitive mutants through chemical mutagenesis, such as with the DNA modifying agent N—methyl—N—nitro—N—nitrosoguanidine; (2) Screening for temperature—sensitive mutants; (3) screening the temperature—sensitive mutants for a block in protein sorting by the use of a construct harboring the staphylococcal enterotoxin B (SEB) gene fused to the cell wall sorting signal of staphylococcal Protein A (SPA), to locate mutants that accumulate a precursor molecule formed by cleavage of an amino—terminal signal peptide but that is not then processed by cleavage of the carboxyl—terminal sorting signal; (4) generation of a S. aureus chromosomal library and complementation of the temperature—sensitive sorting defect; and (5) sequencing and characterization of the S. aureus complementing determinants.

B. Sequence of the Sortase-Transamidase Gene

The above procedure yielded a partial sequence for one of the subunits of the sortase-transamidase including the carboxyl-terminal portion of the gene for the first subunit. This sequence is

GATCCTAAACTGAAAGAAATATATCAAATAGTACTTGAATCTCAAATGAA AGCAATTAATGAGATTAGACCTGGCATGACTGGCAGAAGCTGATGCCA TTTCAAGAAACTATTTAGAGTCAAAAGGGTATGGAAAAGAATTTGGACAT TCACTAGGACATGGTATTGGTTTAGAAATCCATGAAGGGCCAATGCTGGC TCGTACGATACAAGATAAACTTCAAGTTAACAACTGTGTTACAGTAGAAC CTGGTGTTTATAGAAGGTTTGGGCGGTATAAGAATAGAAGATGATATTTT AATTACAGAAAATGGTTGTCAAGTCTTTACTAAATGCACAAAAGACCTTA TAGTTTTAACATAA (SEQ ID NO:28 & 29). The last three nucleotides, TAA, of this sequence are the stop codon.

The above procedure further yielded a sequence for a second subunit of ATGGTCAAAGTAACTGATTATTCAAAATTCAAAATTAGGTAAAGTAGAAAT AGCGCCAGAAGTGCTATCTGTTATTGCAAGTATAGCTACTTCGGAAGTCG AAGGCATCACTGGCCATTTTGCTGAATTAAAAAGAAACAAATTTAGAAAAA GTTAGTCGTAAAAAATTTAAGCCGTGATTTAAAAAATCGAGAGTAAAGAAGA

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TGGCATATATAGATGTATATTGTGCATTAAAACATGGTAATATTTCAAA AACTGCAAACAAAATTCAAACGTCAATTTTTAATTCAAATTTCTAATATGAC AGCGATAGAACCTAAGCAAATTAATATTCACATTACACAAATCGTTATTG AAAAGTAA (SEQ ID NO: 30 & 31) The last three nucleotides of this sequence, TAA, are the stop codon.

Accordingly, within the scope of the present invention is a nucleic acid sequence encoding a substantially purified sortase—transamidase enzyme from a Gram—positive bacterium. The enzyme encoded has at least one subunit with a molecular weight of about 41,000 daltons and catalyzes a reaction that covalently cross—links the carboxyl—terminus of a protein having the sorting signal described above to the peptidoglycan of a gram—positive bacterium. The nucleic acid sequence includes therein the sequence of SEQ ID NO: 28 or a sequence complementary to SEQ ID NO: 28, or the sequence of SEQ ID NO: 30 or a sequence complementary to SEQ ID NO: 30.

Also included within the present invention is a nucleic acid sequence encoding a substantially purified sortase—transamidase enzyme from a Gram—positive bacterium with at least one subunit with a molecular weight of about 41,000 daltons, where the enzyme catalyzes the cross—linking reaction where the nucleic acid sequence hybridizes with at least one of: (1) the sequence of SEQ ID NO: 28; (2) a sequence complementary to SEQ ID NO: 28; (3) the sequence of SEQ ID NO: 30; or (4) a sequence complementary to SEQ ID NO: 30 with no greater than about a 15% mismatch under stringent conditions. Preferably, the degree of mismatch is no greater than about 5%; most preferably the mismatch is no greater than about 2%.

Also within the present invention is a nucleic acid sequence encoding a substantially purified sortase—transamidase enzyme from a Gram—positive bacterium where the enzyme has at least one subunit with a molecular weight of about 41,000 daltons and catalyzes the cross—linking reaction described above involving the sorting signal, where the enzyme includes therein an amino acid sequence selected from the group consisting of: (1) D—P—K—L—K—E—I—Y—Q—I—V—L—E—S—Q—M—K—A—I—N—E—I—R—P—D—M—T—G—A—E—A—D—A—I—S—R—N—Y—L—E—S—K—G—Y—G—K—E—F—G—H—S—L—G—I—G—L—E—I—H—E—G—P—M—L—A—R—T—I—Q—D—K—L—Q—V—N—N—C—V—T—V—E—P—G—V—Y—I—E—G—L—G—I—R—I—E—D—D—I—L—I—T—E—N—G—C—Q—V—F—T—K—C—T—K—D—L—I—V—L—T (SEQ ID NO:2); (2) M—V—K—V—T—D—Y—S—N—S—K—L—G—K—E—I—A—P—E—V—L—S—V—I—A—S—I—A—T—S—E—V—E—G—I—T—G—H—F—A—E—L—K—E—T—N—L—E—K—V—S—R—K—N—L—S—R—D—L—K—I—E—S—K—E—G—I—Y—I—D—V—Y—C—A—L—K—H—G—V—N—I—S—K—T—A—N—K—I—Q—T—S—I—F—N—S—I—S—N—M—T—A—I—E—P—K—O—I—N—I—H—I—T—O—I—V—I—E—K (SEO ID NO:

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31); and (3) sequences incorporating one or more conservative amino acid substitutions in SEQ ID NO:2 or SEQ ID NO: 31 wherein the conservative amino acid substitutions are any of the following: (1) any of isoleucine, leucine and valine for any other of these amino acids; (2) aspartic acid for glutamic acid and vice versa; (3) glutamine for asparagine and vice versa; and (4) serine for threonine and vice versa. Alternative nucleic acid sequences can be determined using the standard genetic code; the alternative codons are readily determinable for each amino acid in this sequence.

Construction of nucleic acid sequences according to the present invention can be accomplished by techniques well known in the art, including solid—phase nucleotide synthesis, the polymerase chain reaction (PCR) technique, reverse transcription of DNA from RNA, the use of DNA polymerases and ligases, and other techniques. If an amino acid sequence is known, the corresponding nucleic acid sequence can be constructed according to the genetic code.

C. Vectors and Host Cells Transformed with Vectors

Another aspect of the invention is a vector comprising a nucleic acid sequence according to the present invention operatively linked to at least one control sequence that controls the expression or regulation of the nucleic acid sequence. Such control sequences are well known in the art and include operators, promoters, enhancers, promoter—proximal elements and replication origins. The techniques of vector construction, including cloning, ligation, gap—filling, the use of the polymerase chain reaction (PCR) procedure, solid—state oligonucleotide synthesis, and other techniques, are all well known in the art and need not be described further here.

Another aspect of the present invention is a host cell transfected with a vector according to the present invention. Among the host cells that can be used are gram-positive bacteria such as *Staphylococcus aureus*.

Transfection, also known as transformation, is done using standard techniques appropriate to the host cell used, particularly *Staphylococcus aureus*. Such techniques are described, for example, in R.P. Novick, "Genetic Systems in Staphylococci," Meth. Enzymol. 204: 587–636 (1991), as well as in O. Schneewind et al., "Sorting of Protein A to the Staphylococcal Cell Wall," Cell 70: 267–281 (1992).

III. SORTASE-TRANSAMIDASE AS A TARGET FOR ANTIBIOTIC ACTION A. A Site for Antibiotic Action

The reaction carried out by the sortase-transamidase of the present invention presents a possible target for a new class of antibiotics to combat medically relevant infections caused by numerous gram-positive organisms. Because this is a

novel site of antibiotic action, these antibiotics have the advantage that resistance by the bacterium has not had a chance to develop.

Such antibiotics can include compounds with structures that mimic the cleavage site, such as compounds with a structure similar to methyl methanethiosulfonate or, more generally, alkyl methanethiosulfonates. The sortasetransamidase of the present invention is believed to be a cysteine protease. Other antibiotics that may inhibit the activity of the sortase-transamidase in the present invention include inhibitors that would be specific for cysteine-modification in a βlactam framework. These inhibitors would have active moieties that would form mixed disulfides with the cysteine sulfhydryl. These active moieties could be 10 derivatives of methanethiosulfonate, such as methanethiosulfonate ethylammonium, methanethiosulfonate ethyltrimethylammonium, or methanethiosulfonate ethylsulfonate (J.A. Javitch et al., "Mapping the Binding Site Crevice of the Dopamine D2 Receptor by the Substituted-Cysteine Accessibility Method," Neuron, 14: 825-831 (1995); M.H. Akabas & A. Karlin, "Identification of Acetylcholine 15 Receptor Channel-Lining Residues in the M1 Segment of the α-Subunit." Biochemistry 34: 12496-12500 (1995)). Similar reagents, such as alkyl alkanethiosulfonates, i.e., methyl methanethiosulfonate, or alkoxycarbonylalkyl disulfides, have been described (D.J. Smith et al., "Simple Alkanethiol Groups for Temporary Blocking of Sulfhydryl Groups of Enzymes," Biochemistry 14: 766-771 20 (1975); W.N. Valentine & D.E. Paglia, "Effect of Chemical Modification of Sulfhydryl Groups of Human Erythrocyte Enzymes," Am. J. Hematol. 11: 111-124 (1981)). Other useful inhibitors involve derivatives of 2-trifluoroacetylaminobenzene sulfonyl fluoride (J.C. Powers, "Proteolytic Enzymes and Their Active-Site-Specific Inhibitors: Role in the Treatment of Disease," in Modification of Proteins), in a β-25 lactam framework, peptidyl aldehydes and nitriles (E. Dufour et al., "Peptide Aldehydes and Nitriles as Transition State Analog Inhibitors of Cysteine Proteases." Biochemistry 34: 9136-9143 (1995); J. O. Westerik & R. Wolfenden, "Aldehydes as Inhibitors of Papain," J. Biol. Chem. 247: 8195-8197 (1972)), peptidyl diazomethyl ketones (L. Björck et al., "Bacterial Growth Blocked by a Synthetic Peptide Based on 30 the Structure of a Human Proteinase Inhibitor," Nature 337: 385-386 (1989)), peptidyl phosphonamidates (P.A. Bartlett & C.K. Marlowe, "Phosphonamidates as Transition-State Analogue Inhibitors of Thermolysin," Biochemistry 22: 4618-4624 (1983)), phosphonate monoesters such as derivatives or analogues of mcarboxyphenyl phenylacetamidomethylphosphonate (R.F. Pratt, "Inhibition of a Class 35 C B-Lactamase by a Specific Phosphonate Monoester," Science 246: 917-919 (1989)), maleimides and their derivatives, including derivatives of such bifunctional

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maleimides as o-phenylenebismaleimide, p-phenylenebismaleimide, mphenylenebismaleimide, 2,3-naphthalenebismaleimide, 1,5naphthalenebismaleimide, and azophenylbismaleimide, as well as monofunctional maleimides and their derivatives (J.V. Moroney et al., "The Distance Between Thiol Groups in the y Subunit of Coupling Factor 1 Influences the Proton Permeability of Thylakoid Membranes," J. Bioenerget. Biomembr. 14: 347–359 (1982)), peptidyl halomethyl ketones (chloromethyl or fluoromethyl ketones), peptidyl sulfonium salts, peptidyl acyloxymethyl ketones, derivatives and analogues of epoxides, such as E-64 (N-[N-(L-trans-carboxyoxiran-2-carbonyl)-L-leucylagmatine), E-64c (a derivative of E-64 in which the agmatine moiety is replaced by an isoamylamine moiety), E-64c ethyl ester, Ep-459 (an analogue of E-64 in which the agmatine moiety is replaced by a 1,4-diaminopropyl moiety), Ep-479 (an analogue of E-64 in which the agmatine moiety is replaced by a 1,7-diheptylamino moiety), Ep-460 (a derivative of Ep-459 in which the terminal amino group is substituted with a Z (benzyloxycarbonyl) group), Ep-174 (a derivative of E-64 in which the agmatine moiety is removed, so that the molecule has a free carboxyl residue from the leucine moiety), Ep-475 (an analogue of E-64 in which the agmatine moiety is replaced with a NH₂-(CH₂)₂-CH-(CH₃)₂ moiety), or Ep-420 (a derivative of E-64 in which the hydroxyl group is benzoylated, forming an ester, and the leucylagmatine mojety is replaced with isoleucyl-O-methyltyrosine), or peptidyl O-acyl hydroxamates (E Shaw, "Cysteinyl Proteases and Their Selective Inactivation), pp 271-347). Other inhibitors are known in the art.

B. Screening Methods

Another aspect of the present invention is a method for screening a compound for anti-sortase-transamidase activity. This is an important aspect of the present invention, because it provides a method for screening for compounds that disrupt the sorting process and thus have potential antibiotic activity against Grampositive bacteria.

In general, this method comprises the steps of: (1) providing an active fraction of sortase—transamidase enzyme; (2) performing an assay for sortase—transamidase activity in the presence and in the absence of the compound being screened; and (3) comparing the activity of the sortase—transamidase enzyme in the presence and in the absence of the compound.

The active fraction of sortase-transamidase enzyme can be a substantially purified sortase-transamidase enzyme preparation according to the

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present invention, but can be a less purified preparation, such as a partially purified particulate preparation as described below.

The enzymatic activity can be measured by the cleavage of a suitable substrate, such as the construct having the Staphylococcal Enterotoxin B (SEB) gene fused to the cell wall sorting signal of Staphylococcal Protein A (SPA). The cleavage can be determined by monitoring the molecular weight of the products by sodium dodecyl sulfate—polyacrylamide gel electrophoresis or by other methods.

One particularly preferred assay for sortase—transamidase activity is the following:

Staphylococcal soluble RNA (sRNA) is prepared from S. aureus by a modification of the technique of Zubay (G. Zubay, <u>J. Mol. Biol.</u> 4: 347–356 (1962)). An overnight culture of S. aureus is diluted 1:10 in TSB and incubated at 37°C for 3 hr. The cells are harvested by centrifugation at 6000 rpm for 15 min.

For every gram of wet cell pellets, 2 ml of 0.01 M magnesium acetate. 0.001 M Tris, pH 7.5 is used to suspend the pellets. The cell pellets are beaten by glass bead beater for 45 minutes in 5 minute intervals. The suspension is centrifuged twice at 2500 rpm for 5 minutes to remove the glass beads, then 0.5 ml phenol is added to the suspension. The suspension is vigorously shaken for 90 minutes at 4°C. and then centrifuged at 18,000 x g for 15 minutes. The nucleic acids in the top layer are precipitated by addition of 0.1 volume of 20% potassium acetate and 2 volumes of ethanol, then stored at 4°C for at least 36 hours. The precipitate is obtained by centrifugation at 5,000 x g for 5 minutes. Cold NaCl (1 ml) is added to the precipitate and stirred at 4°C for 1 hour. The suspension is centrifuged at 15,000 x g for 30 minutes. The sediments are washed with 0.5 ml of cold 1 M NaCl. The supernatants are combined and 2 volumes of ethanol is added to precipitate the tRNA. The precipitate is suspended in 0.1 ml of 0.2 M glycine, pH 10.3 and incubated for 3 hr at 37°C. This suspension is then made 0.4 M in NaCl and the RNA is precipitated by addition of 2 volumes of ethanol. The precipitate is dissolved in 0.7 ml of 0.3 M sodium acetate, pH 7.0. To this is slowly added 0.5 volume of isopropyl alcohol, with stirring. The precipitate is removed by centrifugation at 8,000 x g for 5 min. This precipitate is redissolved in 0.35 ml of 0.3 M sodium acetate, pH 7.0. To this is added 0.5 volume of isopropyl alcohol, using the same procedure as above. The precipitate is also removed by centrifugation. The combined supernatants from the two centrifugations are treated further with 0.37 ml of isopropyl alcohol. The resulting precipitate is dissolved in 75 µl of water and dialyzed against water overnight at 4°C. This sRNA is used in the sortase-transamidase assay.

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Particulate sortase—transamidase enzyme is prepared for use in the assay by a modification of the procedure of Chatterjee & Park (A.N. Chatterjee &-J.T. Park, Proc. Natl. Acad. Sci. USA 51: 9–16 (1964)). An overnight culture of S. aureus OS2 is diluted 1:50 in TSB and incubated at 37°C for 3 hr. Cells are harvested by centrifugation at 6000 rpm for 15 minutes, and washed twice with ice—cold water. The cells are disrupted by shaking 7 ml of 1 3% suspension of cells in 0.05 M Tris—HCl buffer, pH 7.5, 0.1 mM MgCl₂, and 1 mM 2—mercaptoethanol with an equal volume of glass beads for 10–15 minutes in a beater. The glass beads are removed by centrifugation at 2000 rpm for 5 minutes. The crude extract is then centrifuged at 15,000 x g for 5 minutes. The supernatant is centrifuged again at 100,000 x g for 30 minutes. The light yellow translucent pellet is resuspended in 2 to 4 ml of 0.02 M Tris—HCl buffer, pH 7.5, containing 0.1 mM MgCl₂ and 1 mM 2—mercaptoethanol. This suspension represents the crude particulate enzyme and is used in the reaction mixture below.

The supernatant from centrifugation at 100,000 x g is passed through gel filtration using a Sephadex® G-25 agarose column (Pharmacia) to remove endogenous substrates. This supernatant is also used in the reaction mixture.

The complete reaction mixture contains in a final volume of 30 μ l (M. Matsuhashi et al., Proc. Natl. Acad. Sci. USA 54: 587–594 (1965)): 3 μ mol of Tris-HCl, pH 7.8; 0.1 μ mol of MgCl₂; 1.3 μ mol of KCl; 2.7 nmol of [³H] glycine (200 μ Ci/ μ mol); 2 nmol of UDP-M-pentapeptide; 5 nmol of UDP-N-acetylglucosamine; 0.2 μ mol of ATP; 0.05 μ mol of potassium phosphoenolpyruvate; 2.05 μ g of chloramphenicol; 5 μ g of pyruvate kinase; 0.025 μ mol of 2-mercaptoethanol; 50 μ g of staphylococcal sRNA prepared as above; 4 μ g (as protein) of supernatant as prepared above; 271 μ g of particulate enzyme prepared as above; and 8 nmol of a synthesized soluble peptide (HHHHHHHAQALEPTGEENPF) (SEQ ID NO: 32) as a substrate.

The mixture is incubated at 20°C for 60 minutes. The mixture is then heated at 100°C for 1 minute. The mixture is diluted to 1 ml and precipitated with 50 µl nickel resin, and washed with wash buffer (1% Triton X-100, 0.1% sodium dodecyl sulfate, 50 mM Tris, pH 7.5). The nickel resin beads are counted in a scintillation counter to determine ³H bound to the beads.

The effectiveness of the compound being screened to inhibit the activity of the sortase—transamidase enzyme can be determined by adding it to the assay mixture in a predetermined concentration and determining the resulting degree of inhibition of enzyme activity that results. Typically, a dose—response curve is generated using a range of concentrations of the compound being screened.

The particulate enzyme preparation of sortase—transamidase employed in this protocol can be replaced with any other sortase—transamidase preparation, purified or crude, staphylococcal, recombinant, or from any other source from any other Gram—positive bacterium as described above.

The soluble peptide is captured in this embodiment by its affinity for nickel resin as a result of the six histidine residues. More than six histidine residues can be used in the peptide. As an alternative, the soluble peptide can be captured by an affinity resulting from other interactions, such as streptavidin—biotin, glutathione S—transferase—glutathione, maltose binding protein—amylose, and the like, by replacing the six histidine residues with the amino acid sequence that constitutes the binding site in the peptide and employing the appropriate solid phase affinity resin containing the binding partner. Suitable peptides can be prepared by solid phase peptide synthesis using techniques well known in the art, such as those described in M. Bodanszky, "Peptide Chemistry: A Practical Textbook" (2d ed., Springer-Verlag, Berlin, 1993). For example, if the glutathione S—transferase—glutathione interaction is used, the active site of glutathione S—transferase (D.B. Smith & K.S. Johnson, "Single—Step Purification of Polypeptides Expressed in *Escherichia coli* as Fusions with Glutathione S—Transferase," Gene 67: 31–40 (1988)) can be substituted for the six histidine residues, and glutathione can be bound to the solid support.

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IV. <u>USE OF SORTASE-TRANSAMIDASE FOR PROTEIN AND PEPTIDE</u> <u>DISPLAY</u>

A. Methods for Protein and Peptide Display

The sortase-transamidase enzyme of the present invention can also be used in a method of displaying a polypeptide on the surface of a gram-positive bacterium.

In general, a first embodiment of this method comprises the steps of: (1) expressing a polypeptide having a sorting signal at its carboxyl-terminal end as described above; (2) forming a reaction mixture including: (i) the expressed polypeptide; (ii) a substantially purified sortase—transamidase enzyme; and (iii) a Gram—positive bacterium having a peptidoglycan to which the sortase—transamidase can link the polypeptide; and (3) allowing the sortase—transamidase to catalyze a reaction that cleaves the polypeptide within the LPX₃X₄G motif of the sorting signal and covalently cross—links the amino—terminal portion of the cleaved polypeptide to the peptidoglycan to display the polypeptide on the surface of the Gram—positive bacterium.

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In this method, the polypeptide having the sorting signal at its carboxy-terminal end need not be expressed in a Gram-positive bacterium; it can be expressed in another bacterial system such as *Escherichia coli* or *Salmonella typhimurium*, or in a eukaryotic expression system.

The other method for protein targeting and display relies on direct expression of the chimeric protein in a Gram-positive bacterium and the action of the sortase-transamidase on the expressed protein. In general, such a method comprises the steps of: (1) cloning a nucleic acid segment encoding a chimeric protein into a Gram-positive bacterium to generate a cloned chimeric protein including therein a carboxyl-terminal sorting signal as described above, the chimeric protein including the polypeptide to be displayed; (2) growing the bacterium into which the nucleic acid segment has been cloned to express the cloned chimeric protein to generate a chimeric protein including therein a carboxyl-terminal sorting signal; and (3) covalent binding of the chimeric protein to the cell wall by the enzymatic action of the sortase-transamidase involving cleavage of the chimeric protein within the LPX₃X₄G motif so that the protein is displayed on the surface of the gram-positive bacterium in such a way that the protein is accessible to a ligand.

Typically, the Gram-positive bacterium is a species of Staphylococcus. A particularly preferred species of Staphylococcus is Staphylococcus aureus.

However, other Gram-positive bacteria such as *Streptococcus* pyogenes, other *Streptococcus* species, and Gram-positive bacteria of other genera can also be used.

Cloning the nucleic acid segment encoding the chimeric protein into the Gram-positive bacterium is performed by standard methods. In general, such cloning involves: (1) isolation of a nucleic acid segment encoding the protein to be sorted and covalently linked to the cell wall; (2) joining the nucleic acid segment to the sorting signal; (3) cloning by insertion into a vector compatible with the Gram-positive bacterium in which expression is to take place; and (4) incorporation of the vector including the new chimeric nucleic acid segment into the bacterium.

Typically, the nucleic acid segment encoding the protein to be sorted is DNA; however, the use of RNA in certain cloning steps is within the scope of the present invention.

When dealing with genes from eukaryotic organisms, it is preferred to use cDNA, because the natural gene typically contains intervening sequences or introns that are not translated. Alternatively, if the amino acid sequence is known, a synthetic gene encoding the protein to be sorted can be constructed by standard solid-phase oligodeoxyribonucleotide synthesis methods, such as the phosphotriester or

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phosphite triester methods. The sequence of the synthetic gene is determined by the genetic code, by which each naturally occurring amino acid is specified by one or more codons. Additionally, if a portion of the protein sequence is known, but the gene or messenger RNA has not been isolated, the amino acid sequence can be used to construct a degenerate set of probes according to the known degeneracy of the genetic code. General aspects of cloning are described, for example, in J. Sambrook et al., "Molecular Cloning: A Laboratory Manual" (2d ed., Cold Spring Harbor Laboratory Press, Cold Spring Harbor, New York, 1989); in B. Perbal, "A Practical Guide to Molecular Cloning" (2d ed., John Wiley & Sons, New York 1988), in S.L. Berger & A.R. Kimmel, "Guide to Molecular Cloning Techniques" (Methods in Enzymology, vol. 152, Academic Press, Inc., San Diego, 1987), and in D.V. Goeddel, ed., "Gene Expression Technology" (Methods in Enzymology, vol. 185, Academic Press, Inc., San Diego, 1991).

Once isolated, DNA encoding the protein to be sorted is then joined to the sorting signal. This is typically accomplished through ligation, such as using *Escherichia coli* or bacteriophage T4 ligase. Conditions for the use of these enzymes are well known and are described, for example, in the above general references.

The ligation is done in such a way so that the protein to be sorted and the sorting signal are joined in a single contiguous reading frame so that a single protein is produced. This may, in some cases, involve addition or deletion of bases of the cloned DNA segment to maintain a single reading frame. This can be done by using standard techniques.

Cloning is typically performed by inserting the cloned DNA into a vector containing control elements to allow expression of the cloned DNA. The vector is then incorporated into the bacterium in which expression is to occur, using standard techniques of transformation or other techniques for introducing nucleic acids into bacteria.

One suitable cloning system for *S. aureus* places the cloned gene under the control of the BlaZRI regulon (P.Z. Wang et al., <u>Nucl. Acids Res.</u> 19:4000 (1991)). Vectors and other cloning techniques for use in *Staphylococcus aureus* are described in B. Nilsson & L. Abrahmsen, "Fusion to Staphylococcal Protein A," in <u>Gene Expression Technology</u>, <u>supra</u>, p.144–161.

If the chimeric protein is cloned under control of the BlaZRI regulon, expression can be induced by the addition of the β -lactam antibiotic methicillin.

Another aspect of the present invention is a polypeptide displayed on the surface of a Gram-positive bacterium by covalent linkage of an amino-acid sequence of LPX₃X₄ derived from cleavage of an LPX₃X₄G motif, as described above.

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Yet another aspect of the present invention is a covalent complex comprising: (1) the displayed polypeptide; and (2) an antigen or hapten covalently cross-linked to the polypeptide.

B. Screening Methods

These polypeptides associated with the cell surfaces of Gram-positive bacteria can be used in various ways for screening. For example, samples of expressed proteins from an expression library containing expressed proteins on the surfaces of the cells can be used to screen for clones that express a particular desired protein when a labeled antibody or other labeled specific binding partner for that protein is available.

These methods are based on the methods for protein targeting and display described above.

A first embodiment of such a method comprises: (1) expressing a cloned polypeptide as a chimeric protein having a sorting signal at its carboxy—terminal end as described above; (2) forming a reaction mixture including: (i) the expressed chimeric protein; (ii) a substantially purified sortase—transamidase enzyme; and (iii) a Gram—positive bacterium having a peptidoglycan to which the sortase—transamidase can link the polypeptide through the sorting signal; (3) binding of the chimeric protein covalently to the cell wall by the enzymatic action of a sortase—transamidase expressed by the Gram—positive bacterium involving cleavage of the chimeric protein within the LPX₃X₄G motif so that the polypeptide is displayed on the surface of the Gram—positive bacterium in such a way that the polypeptide is accessible to a ligand; and (4) reacting the displayed polypeptide with a labeled specific binding partner to screen the chimeric protein for reactivity with the labeled specific binding partner.

The nucleic acid segment encoding the chimeric protein is formed by methods well known in the art and can include a spacer.

In the last step, the cells are merely exposed to the labeled antibody or other labeled specific binding partner, unreacted antibodies removed as by a wash, and label associated with the cells detected by conventional techniques such as fluorescence, chemiluminescence, or autoradiography.

A second embodiment of this method employs expression in a Gram-positive bacterium that also produces a sortase-transamidase enzyme. This method comprises: (1) cloning a nucleic acid segment encoding a chimeric protein into a Gram-positive bacterium to generate a cloned chimeric protein including therein a carboxyl-terminal sorting signal as described above, the chimeric protein including

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the polypeptide whose expression is to be screened; (2) growing the bacterium into which the nucleic acid segment has been cloned to express the cloned chimeric protein to generate a chimeric protein including therein a carboxyl-terminal sorting signal; (3) binding the polypeptide covalently to the cell wall by the enzymatic action of a sortase-transamidase expressed by the Gram-positive bacterium involving cleavage of the chimeric protein within the LPX₃X₄G motif so that the polypeptide is displayed on the surface of the Gram-positive bacterium in such a way that the polypeptide is accessible to a ligand; and (4) reacting the displayed polypeptide with a labeled specific binding partner to screen the chimeric protein for reactivity with the labeled specific binding partner.

V. <u>USE OF SORTED MOLECULES FOR DIAGNOSIS AND TREATMENT OF</u> BACTERIAL INFECTIONS

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Sorted molecules can also be used for the diagnosis and treatment of bacterial infections caused by Gram-positive bacteria. Antibiotic molecules or fluorescent or any other diagnostic molecules can be chemically linked to a sorted peptide segment, which may include a spacer as described above, and then can be injected into animals or humans. These molecules are then sorted by the sortase-transamidase so that they are covalently linked to the cell wall of the bacteria.

In general, these methods comprise: (1) conjugating an antibiotic or a detection reagent to a protein including therein a carboxyl-terminal sorting signal to produce a conjugate; and (2) introducing the conjugate to an organism infected with a Gram-positive bacterium in order to cause the conjugate to be sorted and covalently cross-linked to the cell walls of the bacterium in order to treat or diagnose the infection.

The antibiotic used can be, but is not limited to, a penicillin, ampicillin, vancomycin, gentamicin, streptomycin, a cephalosporin, amikacin, kanamycin, neomycin, paromomycin, tobramycin, ciprofloxacin, clindamycin, rifampin, chloramphenicol, or norfloxacin, or a derivative of these antibiotics.

The detection reagent is typically an antibody or other specific binding partner labeled with a detectable label, such as a radiolabel. Such methods are well known in the art and need not be described further here.

Accordingly, another aspect of the present invention is a conjugate comprising an antibiotic or a detection reagent covalently conjugated to a protein including therein a carboxyl-terminal sorting signal as described above to produce a conjugate.

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Yet another aspect of the present invention is a composition comprising the conjugate and a pharmaceutically acceptable carrier.

In this context, the conjugates can be administered using conventional modes of administration, including, but not limited to, intravenous, intraperitoneal, oral, or intralymphatic. Other routes of administration can alternatively be used. Oral or intraperitoneal administration is generally preferred. The composition can be administered in a variety of dosage forms, which include, but are not limited to, liquid solutions or suspensions, tablets, pills, powders, suppositories, polymeric microcapsules or microvesicles, liposomes, and injectable or infusible solutions. The preferred form depends on the mode of administration and the quantity administered.

The compositions for administration preferably also include conventional pharmaceutically acceptable carriers and adjuvants known in the art such as human serum albumin, ion exchangers, alumina, lecithin, buffered substances such as phosphate, glycine, sorbic acid, potassium sorbate, and salts or electrolytes such as protamine sulfate. The most effective mode of administration and dosage regimen for the conjugates as used in the methods in the present invention depend on the severity and course of the disease, the patient's health, the response to treatment, the particular strain of bacteria infecting the patient, other drugs being administered and the development of resistance to them, the accessibility of the site of infection to blood flow, pharmacokinetic considerations such as the condition of the patient's liver and/or kidneys that can affect the metabolism and/or excretion of the administered conjugates, and the judgment of the treating physician. According, the dosages should be titrated to the individual patient.

VI. USE OF SORTED POLYPEPTIDES FOR PRODUCTION OF VACCINES

Additionally, the sorted polypeptides covalently crosslinked to the cell walls of Gram-positive bacteria according to the present invention have a number of uses. One use is use in the production of vaccines that can be used to generate immunity against infectious diseases affecting mammals, including both human and non-human mammals, such as cattle, sheep, and goats, as well as other animals such as poultry and fish. This invention is of special importance to mammals. The usefulness of these complexes for vaccine production lies in the fact that the proteins are on the surface of the cell wall and are accessible to the medium surrounding the bacterial cells, so that the antigenic part of the chimeric protein is accessible to the antigen processing system. It is well known that presenting antigens in particulate form greatly enhances the immune response. In effect, bacteria containing antigenic peptides on the surfaces linked to the bacteria by these covalent interactions function

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as natural adjuvants. Here follows a representative list of typical microorganisms that express polypeptide antigens against which useful antibodies can be prepared by the methods of the present invention:

- (1) Fungi: Candida albicans, Aspergillus fumigatus, Histoplasma capsulatum (all cause disseminating disease), Microsporum canis (animal ringworm).
- (2) Parasitic protozoa: (1) *Plasmodium falciparum* (malaria), *Trypanosoma cruzei* (sleeping sickness).
- (3) Spirochetes: (1) Borrelia bergdorferi (Lyme disease), Treponema pallidum (syphilis), Borrelia recurrentis (relapsing fever), Leptospira icterohaemorrhagiae (leptospirosis).
- (4) Bacteria: Neisseria gonorrhoeae (gonorrhea), Staphylococcus aureus (endocarditis), Streptococcus pyogenes (rheumatic fever), Salmonella typhosa (salmonellosis), Hemophilus influenzae (influenza), Bordetella pertussis (whooping cough), Actinomyces israelii (actinomycosis), Streptococcus mutans (dental caries), Streptococcus equi (strangles in horses), Streptococcus agalactiae (bovine mastitis), Streptococcus anginosus (canine genital infections).
- (5) Viruses: Human immunodeficiency virus (HIV), poliovirus, influenza virus, rabies virus, herpes virus, foot and mouth disease virus, psittacosis virus, paramyxovirus, myxovirus, coronavirus.

Typically, the resulting immunological response occurs by both humoral and cell-mediated pathways. One possible immunological response is the production of antibodies, thereby providing protection against infection by the pathogen.

This method is not limited to protein antigens. As discussed below, non-protein antigens or haptens can be covalently linked to the C-terminal cell-wall targeting segment, which can be produced as an independently expressed polypeptide, either alone, or with a spacer at its amino-terminal end. If a spacer at the amino-terminal end is used, typically the spacer will have a conformation allowing the efficient interaction of the non-protein antigen or hapten with the immune system, most typically a random coil or α-helical form. The spacer can be of any suitable length; typically, it is in the range of about 5 to about 30 amino acids; most typically, about 10 to about 20 amino acids. In this version of the embodiment, the independently expressed polypeptide, once expressed, can then be covalently linked to the hapten or non-protein antigen. Typical non-protein antigens or haptens include drugs, including both drugs of abuse and therapeutic drugs, alkaloids, steroids, carbohydrates, aromatic compounds, including many pollutants, and other compounds

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that can be covalently linked to protein and against which an immune response can be raised.

Alternatively, a protein antigen can be covalently linked to the independently expressed cell-wall targeting segment or a cell-wall targeting segment including a spacer.

Many methods for covalent linkage of both protein and non-protein compounds to proteins are well known in the art and are described, for example, in P. Tijssen, "Practice and Theory of Enzyme Immunoassays" (Elsevier, Amsterdam, 1985), pp. 221–295, and in S.S. Wong, "Chemistry of Protein Conjugation and Cross-Linking" (CRC Press, Inc., Boca Raton, FL, 1993).

Many reactive groups on both protein and non-protein compounds are available for conjugation.

For example, organic moieties containing carboxyl groups or that can be carboxylated can be conjugated to proteins via the mixed anhydride method, the carbodiimide method, using dicyclohexylcarbodiimide, and the N-hydroxysuccinimide ester method.

If the organic moiety contains amino groups or reducible nitro groups or can be substituted with such groups, conjugation can be achieved by one of several techniques. Aromatic amines can be converted to diazonium salts by the slow addition of nitrous acid and then reacted with proteins at a pH of about 9. If the organic moiety contains aliphatic amines, such groups can be conjugated to proteins by various methods, including carbodiimide, tolylene–2,4–diisocyanate, or malemide compounds, particularly the N-hydroxysuccinimide esters of malemide derivatives. An example of such a compound is 4–(N-maleimidomethyl)–cyclohexane–1–carboxylic acid. Another example is m-maleimidobenzoyl-N-hydroxysuccinimide ester. Still another reagent that can be used is N-succinimidyl–3–(2–pyridyldithio) propionate. Also, bifunctional esters, such as dimethylpimelimidate, dimethyladipimidate, or dimethylsuberimidate, can be used to couple amino–group–containing moieties to proteins.

Additionally, aliphatic amines can also be converted to aromatic amines by reaction with \underline{p} -nitrobenzoylchloride and subsequent reduction to a \underline{p} -aminobenzoylamide, which can then be coupled to proteins after diazotization.

Organic moieties containing hydroxyl groups can be cross—linked by a number of indirect procedures. For example, the conversion of an alcohol moiety to the half ester of succinic acid (hemisuccinate) introduces a carboxyl group available for conjugation. The bifunctional reagent sebacoyldichloride converts alcohol to acid chloride which, at pH 8.5, reacts readily with proteins. Hydroxyl—containing organic

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moieties can also be conjugated through the highly reactive chlorocarbonates, prepared with an equal molar amount of phosgene.

For organic moieties containing ketones or aldehydes, such carbonyl-containing groups can be derivatized into carboxyl groups through the formation of O-(carboxymethyl) oximes. Ketone groups can also be derivatized with p-hydrazinobenzoic acid to produce carboxyl groups that can be conjugated to the specific binding partner as described above. Organic moieties containing aldehyde groups can be directly conjugated through the formation of Schiff bases which are then stabilized by a reduction with sodium borohydride.

One particularly useful cross-linking agent for hydroxyl-containing organic moieties is a photosensitive noncleavable heterobifunctional cross-linking reagent, sulfosuccinimidyl 6-[4'-azido-2'-nitrophenylamino] hexanoate. Other similar reagents are described in S.S. Wong, "Chemistry of Protein Conjugation and Cross-Linking," supra.

Other cross-linking reagents can be used that introduce spacers between the organic moiety and the specific binding partner.

These methods need not be described further here.

VII. <u>PRODUCTION OF SUBSTANTIALLY PURIFIED SORTASE</u> <u>TRANSAMIDASE ENZYME</u>

Another aspect of the present invention is methods for the production of substantially purified sortase—transamidase enzyme.

A. Methods Involving Expression of Cloned Gene

One method for the production of substantially purified sortase—transamidase enzyme involves the expression of the cloned gene. The isolation of the nucleic acid segment or segments encoding the sortase—transamidase enzyme is described above; these nucleic acid segment or segments are then incorporated into a vector and then use to transform a host in which the enzyme can be expressed. In one alternative, the host is a Gram—positive bacterium.

The next step in this alternative is expression in a Gram-positive bacterium to generate the cloned sortase-transamidase enzyme. Expression is typically under the control of various control elements associated with the vector incorporating the DNA encoding the sortase-transamidase gene; such elements can include promoters and operators, which can be regulated by proteins such as repressors. The conditions required for expression of cloned proteins in grampositive bacteria, particularly *S. aureus*, are well known in the art and need not be

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further recited here. An example is the induction of expression of lysostaphin under control of the BlaZRI regulon induced by the addition of methicillin.

When expressed in *Staphylococcus aureus*, the chimeric protein is typically first exported with an amino-terminal leader peptide, such as the hydrophobic signal peptide at the amino-terminal region of the cloned lysostaphin of Recsei et al. (P. Recsei et al., "Cloning, Sequence, and Expression of the Lysostaphin Gene from *Staphylococcus simulans*," <u>Proc. Natl. Acad. Sci. USA</u> 84:1127-1131 (1987)).

Alternatively, the cloned nucleic acid segment encoding the sortase—transamidase enzyme can be inserted in a vector that contains sequences allowing expression of the sortase—transamidase in another organism, such as *E. coli* or *S. typhimurium*. A suitable host organism can then be transformed or transfected with the vector containing the cloned nucleic acid segment. Expression is then performed in that host organism.

The expressed enzyme is then purified using standard techniques. Techniques for the purification of cloned proteins are well known in the art and need not be detailed further here. One particularly suitable method of purification is affinity chromatography employing an immobilized antibody to sortase. Other protein purification methods include chromatography on ion—exchange resins, gel electrophoresis, isoelectric focusing, and gel filtration, among others.

One particularly useful form of affinity chromatography for purification of cloned proteins, such as sortase—transamidase, as well as other proteins, such as glutathione S—transferase and thioredoxin, that have been extended with carboxyl—terminal histidine residues, is chromatography on a nickel—sepharose column. This allows the purification of a sortase—transamidase enzyme extended at its carboxyl terminus with a sufficient number of histidine residues to allow specific binding of the protein molecule to the nickel—sepharose column through the histidine residues. The bound protein is then eluted with imidazole. Typically, six or more histidine residues are added; preferably, six histidine residues are added. One way of adding the histidine residues to a cloned protein, such the sortase—transamidase, is through PCR with a primer that includes nucleotides encoding the histidine residues. The histidine codons are CAU and CAC expressed as RNA, which are CAT and CAC as DNA. Amplification of the cloned DNA with appropriate primers will add the histidine residues to yield a new nucleic acid segment, which can be recloned into an appropriate host for expression of the enzyme extended with the histidine residues.

B. Other Methods

Alternatively, the sortase-transamidase can be purified from Gram-positive bacteria by standard methods, including precipitation with reagents such as ammonium sulfate or protamine sulfate, ion-exchange chromatography, gel filtration chromatography, affinity chromatography, isoelectric focusing, and gel electrophoresis, as well as other methods known in the art.

Because the sortase—transamidase is a cysteine protease, one particularly useful method of purification involves covalent chromatography by thioldisulfide interchange, using a two-protonic-state gel containing a 2-mercaptopyridine leaving group, such as Sepharose 2B-glutathione 2-pyridyl disulfide or Sepharose 6B-hydroxypropyl 2-pyridyl disulfide. Such covalent chromatographic techniques are described in K. Brocklehurst et al., "Cysteine Proteases," in New Comprehensive Biochemistry, Volume 16: Hydrolytic Enzymes (A. Neuberger & K. Brocklehurst, eds., Elsevier, New York, 1987), ch. 2, pp. 39-158.

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VIII. FURTHER APPLICATIONS OF SORTASE-TRANSAMIDASE

A. <u>Production of Antibodies</u>

Antibodies can be prepared to the substantially purified sortase—transamidase of the present invention, whether the sortase—transamidase is purified from bacteria or produced from recombinant bacteria as a result of gene cloning procedures. Because the substantially purified enzyme according to the present invention is a protein, it is an effective antigen, and antibodies can be made by well—understood methods such as those disclosed in E. Harlow & D. Lane, "Antibodies: A Laboratory Manual" (Cold Spring Harbor Laboratory, 1988). In general, antibody preparation involves immunizing an antibody—producing animal with the protein, with or without an adjuvant such as Freund's complete or incomplete adjuvant, and purification of the antibody produced. The resulting polyclonal antibody can be purified by techniques such as affinity chromatography.

Once the polyclonal antibodies are prepared, monoclonal antibodies can be prepared by standard procedures, such as those described in Chapter 6 of Harlow & Lane, supra.

B. Derivatives for Affinity Chromatography

Another aspect of the present invention is derivatives of the cloned,
substantially purified sortase-transamidase of the present invention extended at its
carboxyl terminus with a sufficient number of histidine residues to allow specific
binding of the protein molecule to a nickel-sepharose column through the histidine

residues. Typically, six or more histidine residues are added; preferably, six histidine residues are added.

The histidine residues can be added to the carboxyl terminus through PCR cloning as described above.

This invention is further described by means of the following example. This Example is for illustrative purposes only, and are not to be construed as limiting the scope of the invention in any manner.

EXAMPLE

Identification of Sortase-Transamidase

Generation of ts Mutants through Chemical Mutagenesis

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To create random mutations in the chromosome, Staphylococcus aureus strain OS2 (RN4220 erm spa-) was mutagenized by exposure to the DNA-modifying agent N-methyl-N-nitro-N-nitrosoguanidine. Cultures were incubated with the mutagen for varying periods of time, then placed on TSB agar plates to measure viability. Cultures were subsequently plated on TSB+ rifampicin (10 μg/ml) to determine the mutation frequency based on resistance to the single target site antibiotic. Once a maximum mutation frequency was reached, cell cultures were exposed to two successive rounds of a penicillin selection (5 μg/ml at 42°C) to enrich for mutants that had a growth effect by lysis of the cells growing at this temperature. Mutants were screened for growth at 30°C and 42°C by streaking individual colonies on TSB agar plates at the permissive and non-permissive temperatures, respectively. These colonies that demonstrated a growth defect at the non-permissive temperature were rechecked at 42°C, and subsequently frozen at -80°C in a 5% BSA, 5% monosodium glutamate (MSG) solution. In this manner, a collection of temperature-sensitive mutants was assembled.

Transformation and Screening of ts Mutants

In order to isolate mutants that demonstrated a defect in the surface display of protein, it was necessary to develop a screening process to locate these strains. Previous studies that had indicated a typical secretory—dependent process, in conjunction with known C—terminal cleavage of translocated proteins, were used to elucidate a selection scheme to isolate the desirable mutants. The construct harboring the staphylococcal enterotoxin B (SEB) gene fused to the cell wall sorting signal of staphylococcal protein A (SPA) was used in this assay. This reporter molecule has been shown to be properly processed not only by the secretory machinery and through signal peptidase cleavage of an N—terminal secretion signal, but also to be correctly

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sorted after secretion to its peptidoglycan substrate. Mutant cells were made competent using 0.5 M sucrose washes, and were transformed with the staphylococcal/Escherichia coli shuttle vector pOS 1 containing the SEB-SPA reporter through electroporation. Transformants were selected by virtue of their resistance to chloramphenicol as encoded by the plasmid. The cells were then screened for properties indicative of a defect in the processing of precursor molecules to a fully matured and anchored surface protein.

Cultured cells were induced at 42°C, and then pulsed with S-35 ProMix (80% Met, 20% Cys), in order to label all synthesized proteins. Samples were precipitated through acid treatment, then digested with lysostaphin (100 µg/ml) and subsequently reprecipitated. This was followed by solubilization in hot sodium dodecyl sulfate (4%), and an immunoprecipitation with anti-SEB antibodies. Samples were run on sodium dodecyl sulfate-polyacrylamide gel electrophoresis and finally exposed to phosphorimager evaluation. The resulting banding patterns were analyzed and quantitated.

This procedure yielded three anti-SEB reacting species, termed P1, P2 and M. P1, the largest precursor which migrated to 33 kDa, represents the complete gene product encoded by the SEB-SPA construct, with no modifications. P2, the second precursor, most likely represents the product after cleavage of the aminoterminal signal peptide found in SEB, and thus migrates to 32 kDa. The smallest species, M, is a lysostaphin-solubilized, maturely anchored peptide that has neither the signal peptide nor the remainder of the carboxyl-terminal sorting signal after the cleavage. This band migrates at approximately 29 kDa.

Analysis of these species was conducted through phosphorimager quantitation, and mutants were selected based upon the proposed phenotype that the inhibition in sorting would result in an accumulation of P2 and a reduction in the production of M. Through this process, two mutants, SM317 and SM329, were earmarked for further analysis due to their elevated ratio of P2/M over wild—type.

Both mutants finally demonstrated an accumulation of P2 after a 5 min kinetic analysis, but also clearly showed a decrease in the mutant's ability to degrade the species over time as measured by samples chased with cold methionine. These results were interpreted to mean that the ability of the mutants to process mature cell wall anchored peptides was impaired, quite possibly due to the less efficient activity of the sortase—transamidase enzyme.

Protoplasts were made of these mutants in order to cure them of their plasmids encoding the reporter construct, and subsequently retransformed with the SEB/SPA containing plasmid to once again test for preservation of this phenotype.

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Upon a favorable result, the mutants were prepared for complementation by a S. aureus chromosomal library.

Generation of an S. aureus Chromosomal Library and Complementation of ts Defect

A staphylococcal library was made through a Sau3AI digest of the chromosomal DNA preparation from *S. aureus* strain RN4220. DNA was isolated through a phenol-chloroform extraction from lysed cells, and digested for various times until the correct partial digest pattern was observed. Fragments greater than 2.5 kb were inserted into the BamHI cloning site in the multi-cloning sequence (MCS) of plasmid pC194-MCS. This heterogeneous mixture of plasmids was then transformed into competent OS2 cells. Approximately 15,000 clones were harvested. DNA was prepared and transformed into competent cells made of both mutants, and simultaneously plated at 30°C and 42°C to screen for complementation of the ts mutant phenotype.

Through this process, four chromosomal inserts from each mutant were found to complement the ts phenotype by conferring growth at 42°C. Due to the nature of the mutagenesis, it is at this point necessary to demonstrate definitively that the ts defect is somehow linked to the defect in processing. This is done through illustration that the plasmids harboring the chromosomal inserts not only complement the temperature sensitivity, but also relieve the accumulation of P2 at the expense of M in these mutants. Therefore, the complemented mutants were screened along with the non-complemented versions against wild type OS2.

Screen for Sorting Defect Complementation

This assay was conducted differently for each of the two mutants. SM-317 was screened by the insertion of the SEB/SPA fragment aboard the replication defective pCL 84 vector that possessed integration capability into the chromosome of *S. aureus* cells. The site-specific integration, mediated by the integrase gene supplied in trans by pCL112, disrupts the lipase gene, which can be assayed for by the lack of hydrolysis on egg yolk agar plates. Once successfully integrated, the RN4220 chromosomal fragments that complement the ts mutation can be added to make the cells ready for screening.

SM329 was assayed by another approach. The pC194 plasmid harboring the complementing stretch of DNA was fused to an *E. coli* replicon pHSG399 that contained the SEB/SPA gene. The shuttle vector thus provided both a reporter substrate as well as a complementing activity.

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The result of this screen demonstrated that over several time periods of the pulse-chase, the complementing insert added to the mutant reduced the accretion of P2 from the elevated level characteristic of the mutant to the wild type range in both mutants, the more dramatically in SM-329. This may be due to the fact that the substrate for sorting in SM-317 is found in only one copy per cell, whereas in SM-329, the reporter aboard pC194 is present at approximately 15 copies. Nevertheless, these results indicate that these mutants are in the sortase-transamidase gene and the sequencing of both chromosomal inserts was therefore undertaken. It should be pointed out that the complementing activity of each of the respective mutants was not transferable to the other, neither in terms of temperature sensitivity nor for the processing defect. Also, the four complementing clones isolated from each mutant seemed to behave in an identical manner, and also seemed to possess very similar restriction sites when digested with various specific endonucleases. Therefore, one clone was chosen from each mutant for sequencing.

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Sequencing and Characterization of the S. aureus Complementing Determinants

The chromosomal inserts carrying the sorting defect complementing capabilities were sequenced. This work was done completely by automated sequence analysis using dideoxyribonucleotides. Sequence data was confirmed by duplicate analysis of both strands of DNA. Comparison was done to all known nucleotide and protein sequence was currently found in the GenBank service. The partial, crude sequence of the *S. aureus* gene is shown in Figure 10 (SEQ ID NO: 28 & 29). The partial carboxy-terminal amino acid sequence of the open reading frame generated from the gene sequence of Figure 10 (SEQ ID NO: 28 & 29) is shown in Figure 11 (SEQ ID NO: 2).

Several stretches of high homology were found, to both known and putative proteins of varying function. The first 364 bases of the SM-317 complementing gene insert been identified as encoding a protein that is a homologue of a putative *Bacillus* peptidase in the GCVT-SPOIIIAA intergenic region (GenBank Accession No. 1731048; Y. Kobayashi et al.). The sequence of this putative peptidase is shown in Figure 5 (SEQ ID NO: 3) and its hydrophobicity profile is shown in Figure 6. The hydrophobicity is calculated according to the method of J. Kyte & R.F. Doolittle, "A Simple Method for Displaying the Hydropathic Character of a Protein," J. Mol. Biol. 157: 105-132 (1982). To a lesser degree of homology, the protein encoded by this complementing gene insert is homologous to aminopeptidase P of *Lactococcus lactis* (GenBank Accession No. 1915907; J. Matos). The amino acid sequence of this aminopeptidase is shown in Figure 7 (SEQ ID NO: 4). To a still

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lesser degree of homology, the protein encoded by this complementing gene insert is homologous to the proline dipeptidase of *Lactobacillus delbrueckii lactis* (GenBank Accession No. 1172066; K. Stucky et al., "Cloning and DNA Sequence Analysis of pepQ, a Prolidase Gene from *Lactobacillus delbrueckii* subsp. *lactis* and Partial Characterization of Its Product," Mol. Gen. Genet. 247: 494-500 (1995). The amino acid sequence of this proline dipeptidase is shown in Figure 8 (SEO ID NO: 5).

An additional complementing gene insert, in a vector designated pCOMP1, has also been sequenced. The DNA sequence of this complementing gene insert is shown in Figure 12 (SEQ ID NO: 30), together with the amino acid sequence (SEQ ID NO: 31) of the protein translated from this DNA sequence, and the hydrophobicity profile of the protein translated from the DNA sequence is shown in Figure 13. The amino acid sequence of the protein translated from the sequence of Figure 12 (SEQ ID NOS: 30 & 31) has virtually no homology with the amino acid sequence of the protein shown in Figure 11 (SEQ ID NO: 2). In particular, the amino acid sequence of the protein of Figure 12 (SEQ ID NOS: 30 & 31) has a single cysteine residue.

Although Applicants do not intend to be bound by this theory, the existence of these two complementing inserts that define different polypeptide sequences suggests that the sortase—transamidase enzyme of *S. aureus* is a heterooligomer, with two or more different subunits that have different amino acid sequences. Mutations in at least two different subunits can give rise to the temperature sensitive phenotype and can then be complemented for. Alternatively, the synthesis of the peptidoglycan may require additional enzymes.

Upon completion of the sequencing, and a study of all open reading frames, candidate genes were selected for further analysis. These genes were expressed in the mutants to determine if they complement both the ts and sorting defects. Upon success in this capacity, the genes were disrupted in wild—type S. aureus to determine their essentiality and possible biological roles.

30 Materials and Methods

Mutagenesis of S. aureus Strain OS2. S. aureus strain OS2 (RN4220 erm spa-) was mutagenized by treatment with N-methyl-N-nitro-N-nitrosoguanidine at 2 mg/ml. Culture OD was measured at 660 nm for viability. After 45 min, cultures were spun down at 4,000 rpm for 10 min, and washed with citrate buffer, pH 5.5. Cells were resuspended in citrate buffer to a concentration of 5x10⁸ cells/ml. These cultures were serially diluted in TSB (tryptic soy broth) and

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plated on TSB+ rifampicin (10 μ g/ml) agar plates. Mutation frequencies were determined to be $5x10^5$ mutation/cfu.

Enrichment and Selection of ts Mutants. A 1:100 dilution of cell cultures grown overnight at 30°C was added to TSB, and allowed to grow for 2 hr, at which time penicillin G was added at 5 μ g/ml. Culture viability was measured by taking OD readings at 660 nm at various times until the concentration dropped to a stable point. Cells were washed twice in TSB, and the enrichment was repeated a second time.

After two successive rounds of penicillin selection for growth arrest at 42°C, individual colonies were picked and simultaneously streaked on duplicate TSB plates incubated at 30°C and 42°C. Those colonies that had a growth defect at 42°C were rechecked at this nonpermissive temperature, and stored at -80°C in a 5% BSA 5% MSG solution.

Transformation of Competent Cells. Mutant cells were made competent by diluting overnight cultures 1:10 in TSB, and growing to 0.3 OD at 660 nm. Cells were then spun down at 7,500 rpm for 15 min and resuspended in an equal volume of 0.5 M sucrose. After another pelleting, cells were resuspended in 0.5 volume sucrose, and incubated for 30 min at 4°C. Following another spin, cells were brought up in 0.1 volume sucrose. These competent cells were then transformed with the appropriate plasmids encoding chloramphenicol (10 μ g/ml) resistance by electroporation at 200 ohms, 25 μ F, and 2.5 kV in 0.2 cm cuvettes. Cells were plated on TSB plus chloramphenicol incubated at 30°C. Transformed strains were frozen in BSA/MSG at -80°C.

Pulse-Chase Screen of Mutants. Strains were inoculated in chemically defined media with chloramphenicol and grown overnight at 30°C. Cultures were 25 diluted 1:10 into medium IV (O. Schneewind et al., "Sorting of Protein A to the Staphylococcal Cell Wall," Cell 70: 267-281 (1992)), grown for 3 hr, and then induced at 42°C for 20 min. At this time, cultures were pulse labeled with 50 µCi of S-35 ProMix for 5 min, and then terminated with 5% trichloroacetic acid (TCA). Cells were incubated at 4° for 30 min, centrifuged at 12,500 rpm for 15 min, and the 30 supernatants aspirated. After resuspension in acetone, cells were spun again and aspirated to dryness. At this time, cells were treated with lysostaphin (100 µg/ml) for 30 min or until noticeable clearing, and subjected to another TCA/acetone precipitation. After lysis of cells by boiling for 10 min with 4% SDS in 0.5m Tris. pH 8.0, proteins were immunoprecipitated with anti-SEB for 1 hr and protein A-35 Sepharose beads for another 1 hr. Samples were washed three times in RIPA buffer. pH 8.0 (300 mM NaCl, 2% Triton X-100, 1% deoxycholate, 0.2% SDS), and protein

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was solubilized with SDS-urea sample buffer (50 mM Tris-HCl, pH 6.8, 3% SDS, 5% 2-mercaptoethanol, 3.5 M urea), with boiling for 10 min. Samples were run on SDS-polyacrylamide gel electrophoresis gels and exposed to a phosphorimager screen overnight. Quantitations were done on ImageQuant software.

<u>DNA Sequencing</u>. DNA was sequenced on a Perkin-Elmer automated sequencer after PCR using dye-terminating ready reaction mixed with SS-Taq polymerase. GenBank analysis was done using BLAST software to search the database.

ADVANTAGES OF THE PRESENT INVENTION

In isolating and characterizing the gene for the S. aureus sortase—transamidase enzyme, we have determined the existence of a new site for antibiotic action that can be used to screen new antibiotics active against Gram—positive pathogens, such as Staphylococcus, Actinomyces, Mycobacterium, Streptococcus, Bacillus, and other medically important Gram—positive pathogens increasingly resistant to conventional antibiotics. The availability of substantially purified S. aureus sortase—transamidase enzyme provides a method of screening compounds for inhibition of the enzyme.

The purified sortase—transamidase enzyme of the present invention also yields a method of surface display of peptides and proteins that has advantages over phage display, as well as providing methods for producing vaccines against a large variety of antigens that can be covalently bound to the surfaces of Grampositive bacteria.

Although the present invention has been described with considerable
detail, with reference to certain preferred versions thereof, other versions and
embodiments are possible. Therefore, the scope of the invention is determined by the
following claims.

We claim:

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1. A substantially purified sortase—transamidase enzyme from a

Gram—positive bacterium, the enzyme catalyzing a reaction that covalently cross—
links the carboxyl terminus of a protein having a sorting signal to the peptidoglycan of a Gram—positive bacterium, the sorting signal having a motif of LPX₃X₄G therein, wherein sorting occurs by cleavage between the fourth and fifth residues of the LPX₃X₄G motif.

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2. The substantially purified sortase—transamidase enzyme of claim 1 wherein the Gram—positive bacterium is a species selected from the group consisting of Staphylococcus aureus, S. sobrinus, Enterococcus faecalis, Streptococcus pyogenes, and Listeria monocytogenes.

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- 3. The substantially purified sortase-transamidase enzyme of claim 2 wherein the Gram-positive bacterium is *Staphylococcus aureus*.
- 4. The substantially purified sortase-transamidase enzyme of claim 1 wherein one subunit of the enzyme has a molecular weight of about 41,000 daltons.
 - 5. The substantially purified sortase—transamidase enzyme of claim 4 wherein the sorting signal further comprises:(2) a substantially hydrophobic domain of at least 31 amino acids carboxyl to the motif; and (3) a charged tail region with at least two positively charged residues carboxyl to the substantially hydrophobic domain, at least one of the two positively charged residues being arginine, the two positively charged residues being located at residues 31–33 from the motif, wherein X₃ is any of the twenty naturally—occurring L—amino acids and X₄ is selected from the group consisting of alanine, serine, and threonine.

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6. The enzyme of claim 1 wherein the enzyme includes therein an amino acid sequence selected from the group consisting of :(1) D-P-K-L-K-E-I-Y-Q-I-V-L-E-S-Q-M-K-A-I-N-E-I-R-P-G-M-T-G-A-E-A-D-A-I-S-R-N-Y-L-E-S-K-G-Y-G-K-E-F-G-H-S-L-G-H-G-I-G-L-E-I-H-E-G-P-M-L-A-R-T-I-Q-D-K-L-Q-V-N-N-C-V-T-V-E-P-G-V-Y-I-E-G-L-G-I-R-I-E-D-D-I-L-I-T-E-N-G-C-Q-V-F-T-K-C-T-K-D-L-I-V-L-T (SEQ ID NO: 2); (2) M-V-K-V-T-D-Y-S-N-S-K-L-G-K-E-I-A-P-E-V-L-

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S-V-I-A-S-I-A-T-S-E-V-E-G-I-T-G-H-F-A-E-L-K-E-T-N-L-E-K-V-S-R-K-N-L-S-R-D-L-K-I-E-S-K-E-G-I-Y-I-D-V-Y-C-A-L-K-H-G-V-N-I-S-K-T-A-N-K-I-Q-T-S-I-F-N-S-I-S-N-M-T-A-I-E-P-K-Q-I-N-I-H-I-T-Q-I-V-I-E-K (SEQ ID NO: 31); and (3) sequences incorporating one or more conservative amino acid substitutions in SEQ ID NO: 2 or SEQ ID NO: 31, wherein the conservative amino acid substitutions are any of the following: (1) any of isoleucine, leucine, and valine for any other of these amino acids; (2) aspartic acid for glutamic acid and vice versa; (3) glutamine for asparagine and vice versa; and (4) serine for threonine and vice versa.

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- 7. The enzyme of claim 6 wherein the amino acid sequence is D-P-K-L-K-E-I-Y-Q-I-V-L-E-S-Q-M-K-A-I-N-E-I-R-P-G-M-T-G-A-E-A-D-A-I-S-R-N-Y-L-E-S-K-G-Y-G-K-E-F-G-H-S-L-G-H-G-I-G-L-E-I-H-E-G-P-M-L-A-R-T-I-Q-D-K-L-Q-V-N-N-C-V-T-V-E-P-G-V-Y-I-E-G-L-G-I-R-I-E-D-D-I-L-I-T-E-N-G-C-Q-V-F-T-K-C-T-K-D-L-I-V-L-T (SEQ ID NO: 2).
- 8. The enzyme of claim 6 wherein the amino acid sequence is M-V-K-V-T-D-Y-S-N-S-K-L-G-K-E-I-A-P-E-V-L-S-V-I-A-S-I-A-T-S-E-V-E-G-I-T-G-H-F-A-E-L-K-E-T-N-L-E-K-V-S-R-K-N-L-S-R-D-L-K-I-E-S-K-E-G-I-Y-I-D-V-Y-C-A-L-K-H-G-V-N-I-S-K-T-A-N-K-I-Q-T-S-I-F-N-S-I-S-N-M-T-A-I-E-P-K-Q-I-N-I-H-I-T-Q-I-V-I-E-K (SEQ ID NO: 31).

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- 9. A nucleic acid sequence encoding the enzyme of claim 6.
- 10. A nucleic acid sequence encoding the enzyme of claim 7.
- 11. A nucleic acid sequence encoding the enzyme of claim 8.

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12. A nucleic acid sequence encoding a substantially purified sortase—transamidase enzyme from a Gram—positive bacterium, the enzyme having a subunit with a molecular weight of about 41,000 daltons and catalyzing a reaction that covalently cross—links the carboxyl terminus of a protein having a sorting signal to the peptidoglycan of a Gram—positive bacterium, the sorting signal having: (1) a motif of LPX₃X₄G therein; (2) a substantially hydrophobic domain of at least 31 amino acids carboxyl to the motif; and (3) a charged tail region with at least two positively charged

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residues carboxyl to the substantially hydrophobic domain, at least one of the two positively charged residues being arginine, the two positively charged residues being located at residues 31-33 from the motif, wherein X3 is any of the twenty naturallyoccurring L-amino acids and X4 is selected from the group consisting of alanine, serine, and threonine, and wherein sorting occurs by cleavage between the fourth and 5 fifth residues of the LPX₃X₄G motif, wherein the nucleic acid sequence includes therein a sequence selected from the group consisting of: (1) GATCCTAAACTGAAAGAAATATATCAAATAGTACTTGAATCTCAAATGAA AGCAATTAATGAGATTAGACCTGGCATGACTGGTGCAGAAGCTGATGCCA TTTCAAGAAACTATTTAGAGTCAAAAGGGTATGGAAAAGAATTTGGACAT 10 TCACTAGGACATGGTATTGGTTTAGAAATCCATGAAGGGCCAATGCTGGC TCGTACGATACAAGATAAACTTCAAGTTAACAACTGTGTTACAGTAGAAC CTGGTGTTTATATAGAAGGTTTGGGCGGTATAAGAATAGAAGATGATATT TTAATTACAGAAAATGGTTGTCAAGTCTTTACTAAATGCACAAAAGACCTT ATAGTTTTAACATAA (SEQ ID NO: 28); (2) 15 ATGGTCAAAGTAACTGATTATTCAAAATTCAAAATTAGGTAAAGTAGAAAT AGCGCCAGAAGTGCTATCTGTTATTGCAAGTATAGCTACTTCGGAAGTCG AAGGCATCACTGGCCATTTTGCTGAATTAAAAGAAACAAATTTAGAAAAA GTTAGTCGTAAAAATTTAAGCCGTGATTTAAAAATCGAGAGTAAAGAAGA TGGCATATATAGATGTATATTGTGCATTAAAACATGGTAATATTTCAAA 20 AACTGCAAACAAAATTCAAACGTCAATTTTTAATTCAATTTCTAATATGAC AGCGATAGAACCTAAGCAAATTAATATTCACATTACACAAATCGTTATTG AAAAGTAA (SEQ ID NO: 30); or (3) a sequence complementary to SEO ID NO: 28

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or SEQ ID NO: 30.

transamidase enzyme from a Gram-positive bacterium, the enzyme having a subunit with a molecular weight of about 41,000 daltons and catalyzing a reaction that covalently cross-links the carboxyl terminus of a protein having a sorting signal to the peptidoglycan of a Gram-positive bacterium, the sorting signal having (1) a motif of LPX₃X₄G therein; (2) a substantially hydrophobic domain of at least 31 amino acids carboxyl to the motif; and (3) a charged tail region with at least two positively charged residues carboxyl to the substantially hydrophobic domain, at least one of the two positively charged residues being arginine, the two positively charged residues being located at residues 31-33 from the motif, wherein X₃ is any of the twenty naturally-occurring L-amino acids and X₄ is selected from the group consisting of alanine, serine, and threonine, and wherein sorting occurs by cleavage between the fourth and

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fifth residues of the LPX₃X₄G motif, wherein the nucleic acid sequence hybridizes with a sequence selected from the group consisting of: (1) GATCCTAAACTGAAAGAAATATATCAAATAGTACTTGAATCTCAAATGAA AGCAATTAATGAGATTAGACCTGGCATGACTGGTGCAGAAGCTGATGCCA TTTCAAGAAACTATTTAGAGTCAAAAGGGTATGGAAAAGAATTTGGACAT TCACTAGGACATGGTATTGGTTTAGAAATCCATGAAGGGCCAATGCTGGC TCGTACGATACAAGATAAACTTCAAGTTAACAACTGTGTTACAGTAGAAC CTGGTGTTTATATAGAAGGTTTGGGCGGTATAAGAATAGAAGATGATATT TTAATTACAGAAAATGGTTGTCAAGTCTTTACTAAATGCACAAAAGACCTT ATAGTTTTAACATAA (SEQ ID NO: 28); (2) ATGGTCAAAGTAACTGATTATTCAAAATTCAAAATTAGGTAAAGTAGAAAT AGCGCCAGAAGTGCTATCTGTTATTGCAAGTATAGCTACTTCGGAAGTCG AAGGCATCACTGGCCATTTTGCTGAATTAAAAGAAACAAATTTAGAAAA GTTAGTCGTAAAAATTTAAGCCGTGATTTAAAAATCGAGAGTAAAGAAGA TGGCATATATATAGATGTATATTGTGCATTAAAACATGGTAATATTTCAAA 15 AACTGCAAACAAATTCAAACGTCAATTTTTAATTCAATTTCTAATATGAC AGCGATAGAACCTAAGCAAATTAATATTCACATTACACAAATCGTTATTG AAAAGTAA (SEQ ID NO: 30) or (3) a sequence complementary to SEO ID NO: 28

- 14. The nucleic acid sequence of claim 13 wherein the mismatch is no greater than about 5%.
- 25 15. The nucleic acid sequence of claim 14 wherein the mismatch is no greater than about 2%.

or SEQ ID NO: 30, with no greater than about a 15% mismatch under stringent

- 16. A vector comprising the nucleic acid sequence of claim 9 operatively linked to at least one control sequence that controls the expression or regulation of the nucleic acid sequence.
- 17. A vector comprising the nucleic acid sequence of claim 10 operatively linked to at least one control sequence that controls the expression or regulation of the nucleic acid sequence.

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conditions.

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- 18. A vector comprising the nucleic acid sequence of claim 11 operatively linked to at least one control sequence that controls the expression or regulation of the nucleic acid sequence.
- 5 19. A vector comprising the nucleic acid sequence of claim 12 operatively linked to at least one control sequence that controls the expression or regulation of the nucleic acid sequence.
- 20. A vector comprising the nucleic acid sequence of claim 13 operatively linked to at least one control sequence that controls the expression or regulation of the nucleic acid sequence.
 - 21. A host cell transfected with the vector of claim 16.
 - 22. A host cell transfected with the vector of claim 17.
 - 23. A host cell transfected with the vector of claim 18.
 - 24. A host cell transfected with the vector of claim 19.
 - 25. A host cell transfected with the vector of claim 20.
 - 26. A method for producing a substantially purified sortase-transamidase enzyme comprising the steps of:
 - (a) culturing the host cell of claim 21 under conditions in which the host cell expresses the encoded sortase-transamidase enzyme; and
 - (b) purifying the expressed enzyme to produce substantially purified sortase-transamidase enzyme.
 - 27. A method for producing a substantially purified sortase—transamidase enzyme comprising the steps of:
 - (a) culturing the host cell of claim 22 under conditions in which the host cell expresses the encoded sortase-transamidase enzyme; and
- (b) purifying the expressed enzyme to produce substantially purified sortase-transamidase enzyme.

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- 28. A method for producing a substantially purified sortase-transamidase enzyme comprising the steps of:
- (a) culturing the host cell of claim 23 under conditions in which the host cell expresses the encoded sortase-transamidase enzyme; and
- (b) purifying the expressed enzyme to produce substantially purified sortase-transamidase enzyme.
- 29. A method for producing a substantially purified sortase—transamidase enzyme comprising the steps of:
- (a) culturing the host cell of claim 24 under conditions in which the host cell expresses the encoded sortase-transamidase enzyme; and
- (b) purifying the expressed enzyme to produce substantially purified sortase-transamidase enzyme.
- 30. A method for producing a substantially purified sortase—transamidase enzyme comprising the steps of:
- (a) culturing the host cell of claim 25 under conditions in which the host cell expresses the encoded sortase-transamidase enzyme; and
- (b) purifying the expressed enzyme to produce substantially purified sortase-transamidase enzyme.
 - 31. Substantially purified sortase-transamidase enzyme produced by the process of claim 26.
- 32. Substantially purified sortase—transamidase enzyme produced by the process of claim 27.
 - 33. Substantially purified sortase-transamidase enzyme produced by the process of claim 28.
 - 34. Substantially purified sortase—transamidase enzyme produced by the process of claim 29.
- 35. Substantially purified sortase-transamidase enzyme produced by the process of claim 30.

- 36. A method for screening a compound for anti-sortase-transamidase activity comprising the steps of:
- (a) providing the substantially purified sortase-transamidase enzyme of claim 1;
- (b) performing an assay for sortase-transamidase in the presence and in the absence of the compound; and
- (c) comparing the activity of the sortase-transamidase enzyme in the presence and in the absence of the compound to screen the compound for sortase-transamidase activity.

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- 37. A method for screening a compound for anti-sortase-transamidase activity comprising the steps of:
- (a) providing the substantially purified sortase-transamidase enzyme of claim 3;
- (b) performing an assay for sortase-transamidase in the presence and in the absence of the compound; and
- (c) comparing the activity of the sortase—transamidase enzyme in the presence and in the absence of the compound to screen the compound for sortase—transamidase activity.

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- 38. A method for screening a compound for anti-sortase-transamidase activity comprising the steps of:
- (a) providing the substantially purified sortase-transamidase enzyme of claim 31;
- (b) performing an assay for sortase-transamidase in the presence and in the absence of the compound; and
- (c) comparing the activity of the sortase—transamidase enzyme in the presence and in the absence of the compound to screen the compound for sortase—transamidase activity.

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- 39. A method for screening a compound for anti-sortase-transamidase activity comprising the steps of:
- (a) providing the substantially purified sortase—transamidase enzyme of claim 32;
- 35 (b) performing an assay for sortase-transamidase in the presence and in the absence of the compound; and

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- (c) comparing the activity of the sortase—transamidase enzyme in the presence and in the absence of the compound to screen the compound for sortase—transamidase activity.
- 40. A method for screening a compound for anti-sortase-transamidase activity comprising the steps of:
 - (a) providing the substantially purified sortase-transamidase enzyme of claim 33;
- (b) performing an assay for sortase-transamidase in the presence and in the absence of the compound; and
 - (c) comparing the activity of the sortase—transamidase enzyme in the presence and in the absence of the compound to screen the compound for sortase—transamidase activity.
 - 41. A method for screening a compound for anti-sortase-transamidase activity comprising the steps of:
 - (a) providing the substantially purified sortase-transamidase enzyme of claim 34;
 - (b) performing an assay for sortase-transamidase in the presence and in the absence of the compound; and
 - (c) comparing the activity of the sortase—transamidase enzyme in the presence and in the absence of the compound to screen the compound for sortase transamidase activity.
- 42. A method for screening a compound for anti-sortase-transamidase activity comprising the steps of:
 - (a) providing the substantially purified sortase-transamidase enzyme of claim 35;
- (b) performing an assay for sortase-transamidase in the presence and in the absence of the compound; and
 - (c) comparing the activity of the sortase—transamidase enzyme in the presence and in the absence of the compound to screen the compound for sortase—transamidase activity.

- 43. A method for screening a compound for anti-sortase-transamidase activity comprising the steps of:
- (a) providing an active fraction of sortase-transamidase enzyme from a Gram-positive bacterium;
- (b) performing an assay for sortase-transamidase in the presence and in the absence of the compound; and
- (c) comparing the activity of the sortase—transamidase enzyme in the presence and in the absence of the compound to screen the compound for sortase—transamidase activity.

44. The method of claim 43 wherein the active fraction of sortase—transamidase enzyme is a particulate fraction from *Staphylococcus aureus*.

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- 45. The method of claim 43 wherein the assay for sortase—
 transamidase enzyme is performed by monitoring the capture of a soluble peptide that is a substrate for the enzyme by its interaction with an affinity resin.
 - 46. The method of claim 45 wherein the soluble peptide includes a sequence of at least six histidine residues and the affinity resin contains nickel.
 - 47. The method of claim 45 wherein the soluble peptide includes the active site of glutathione S-transferase and the affinity resin contains glutathione.
- 48. The method of claim 45 wherein the soluble peptide includes the active site of streptavidin and the affinity resin contains biotin.
 - 49. The method of claim 45 wherein the soluble peptide includes the active site of maltose binding protein and the affinity resin contains amylose.
 - 50. An antibody specifically binding the substantially purified sortase—transamidase enzyme of claim 1.
 - 51. An antibody specifically binding the substantially purified sortase—transamidase enzyme of claim 3.
 - 52. An antibody specifically binding the substantially purified sortase-transamidase enzyme of claim 31.

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- 53. An antibody specifically binding the substantially purified sortase-transamidase enzyme of claim 32.
- 5 54. An antibody specifically binding the substantially purified sortase-transamidase enzyme of claim 33.
 - 55. An antibody specifically binding the substantially purified sortase-transamidase enzyme of claim 34.
 - 56. An antibody specifically binding the substantially purified sortase-transamidase enzyme of claim 35.
- 57. A protein molecule comprising the substantially purified sortase—
 transamidase enzyme of claim 1 extended at its carboxyl—terminus with a sufficient number of histidine residues to allow specific binding of the protein molecule to a nickel—sepharose column through the histidine residues added at the carboxyl—terminus.
- 58. A protein molecule comprising the substantially purified sortase—
 transamidase enzyme of claim 3 extended at its carboxyl—terminus with a sufficient
 number of histidine residues to allow specific binding of the protein molecule to a
 nickel—sepharose column through the histidine residues added at the carboxyl—
 terminus.
 - 59. A protein molecule comprising the substantially purified sortase-transamidase enzyme of claim 31 extended at its carboxyl-terminus with a sufficient number of histidine residues to allow specific binding of the protein molecule to a nickel-sepharose column.
 - 60. A protein molecule comprising the substantially purified sortase—transamidase enzyme of claim 32 extended at its carboxyl-terminus with a sufficient number of histidine residues to allow specific binding of the protein molecule to a nickel-sepharose column.

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61. A protein molecule comprising the substantially purified sortase—transamidase enzyme of claim 33 extended at its carboxyl—terminus with a sufficient number of histidine residues to allow specific binding of the protein molecule to a nickel—sepharose column.

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62. A protein molecule comprising the substantially purified sortase—transamidase enzyme of claim 34 extended at its carboxyl—terminus with a sufficient number of histidine residues to allow specific binding of the protein molecule to a nickel—sepharose column.

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63. A protein molecule comprising the substantially purified sortase—transamidase enzyme of claim 35 extended at its carboxyl—terminus with a sufficient number of histidine residues to allow specific binding of the protein molecule to a nickel—sepharose column.

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- 64. A method for displaying a polypeptide on the surface of a Grampositive bacterium comprising the steps of:
- (a) expressing a polypeptide having a sorting signal at its carboxy—terminal end, the sorting signal having: (1) a motif of LPX₃X₄G therein; (2) a substantially hydrophobic domain of at least 31 amino acids carboxyl to the motif; and (3) a charged tail region with at least two positively charged residues carboxyl to the substantially hydrophobic domain, at least one of the two positively charged residues being arginine, the two positively charged residues being located at residues 31–33 from the motif, wherein X₃ is any of the twenty naturally—occurring L—amino acids and X₄ is selected from the group consisting of alanine, serine, and threonine;
- (b) forming a reaction mixture including: (i) the expressed polypeptide; (ii) the substantially purified sortase-transamidase of claim 1; and (iii) a Gram-positive bacterium having a peptidoglycan to which the sortase-transamidase can link the polypeptide; and

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(c) allowing the sortase-transamidase to catalyze a reaction that cleaves the polypeptide within the LPX_3X_4 motif of the sorting signal and covalently cross-links the amino-terminal portion of the cleaved polypeptide to the peptidoglycan to display the polypeptide on the surface of the Gram-positive bacterium.

- 65. A method for displaying a polypeptide on the surface of a Gram-positive bacterium comprising the steps of:
- (a) expressing a polypeptide having a sorting signal at its carboxy—terminal end, the sorting signal having: (1) a motif of LPX₃X₄G therein; (2) a substantially hydrophobic domain of at least 31 amino acids carboxyl to the motif; and (3) a charged tail region with at least two positively charged residues carboxyl to the substantially hydrophobic domain, at least one of the two positively charged residues being arginine, the two positively charged residues being located at residues 31–33 from the motif, wherein X₃ is any of the twenty naturally—occurring L—amino acids and X₄ is selected from the group consisting of alanine, serine, and threonine;
- (b) forming a reaction mixture including: (i) the expressed polypeptide; (ii) the substantially purified sortase-transamidase of claim 3; and (iii) a Gram-positive bacterium having a peptidoglycan to which the sortase-transamidase can link the polypeptide; and
- (c) allowing the sortase—transamidase to catalyze a reaction that cleaves the polypeptide within the LPX₃X₄G motif of the sorting signal and covalently cross—links the amino—terminal portion of the cleaved polypeptide to the peptidoglycan to display the polypeptide on the surface of the Gram—positive bacterium.

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- 66. A method for displaying a polypeptide on the surface of a Gram-positive bacterium comprising the steps of:
- (a) expressing a polypeptide having a sorting signal at its carboxy-terminal end, the sorting signal having: (1) a motif of LPX₃X₄G therein; (2) a substantially hydrophobic domain of at least 31 amino acids carboxyl to the motif; and (3) a charged tail region with at least two positively charged residues carboxyl to the substantially hydrophobic domain, at least one of the two positively charged residues being arginine, the two positively charged residues being alocated at residues 31-33 from the motif, wherein X_3 is any of the twenty naturally-occurring L-amino acids and X_4 is selected from the group consisting of alanine, serine, and threonine:
- (b) forming a reaction mixture including: (i) the expressed polypeptide; (ii) the substantially purified sortase-transamidase enzyme of claim 31; and (iii) a Gram-positive bacterium having a peptidoglycan to which the sortase-transamidase can link the polypeptide; and
- (c) allowing the sortase-transamidase to catalyze a reaction that cleaves the polypeptide within the LPX₃X₄G motif of the sorting signal and covalently cross-links the amino-terminal portion of the cleaved polypeptide to the

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peptidoglycan to display the polypeptide on the surface of the Gram-positive bacterium.

- 67. A method for displaying a polypeptide on the surface of a Gram-5 positive bacterium comprising the steps of:
 - (a) expressing a polypeptide having a sorting signal at its carboxyterminal end, the sorting signal having: (1) a motif of LPX₃X₄G therein; (2) a substantially hydrophobic domain of at least 31 amino acids carboxyl to the motif; and (3) a charged tail region with at least two positively charged residues carboxyl to the substantially hydrophobic domain, at least one of the two positively charged residues being arginine, the two positively charged residues being located at residues 31-33 from the motif, wherein X₃ is any of the twenty naturally-occurring L-amino acids and X₄ is selected from the group consisting of alanine, serine, and threonine:

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- (b) forming a reaction mixture including: (i) the expressed polypeptide; (ii) the substantially purified sortase-transamidase enzyme of claim 32: and (iii) a Gram-positive bacterium having a peptidoglycan to which the sortasetransamidase can link the polypeptide; and
- (c) allowing the sortase-transamidase to catalyze a reaction that cleaves the polypeptide within the LPX₃X₄G motif of the sorting signal and covalently cross-links the amino-terminal portion of the cleaved polypeptide to the peptidoglycan to display the polypeptide on the surface of the Gram-positive bacterium.
- 68. A method for displaying a polypeptide on the surface of a Grampositive bacterium comprising the steps of:
 - (a) expressing a polypeptide having a sorting signal at its carboxyterminal end, the sorting signal having: (1) a motif of LPX₃X₄G therein; (2) a substantially hydrophobic domain of at least 31 amino acids carboxyl to the motif; and (3) a charged tail region with at least two positively charged residues carboxyl to the substantially hydrophobic domain, at least one of the two positively charged residues being arginine, the two positively charged residues being located at residues 31-33 from the motif, wherein X₃ is any of the twenty naturally-occurring L-amino acids and X₄ is selected from the group consisting of alanine, serine, and threonine:
- (b) forming a reaction mixture including: (i) the expressed polypeptide; (ii) the substantially purified sortase-transamidase enzyme of claim 33; 35 and (iii) a Gram-positive bacterium having a peptidoglycan to which the sortasetransamidase can link the polypeptide; and

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- (c) allowing the sortase-transamidase to catalyze a reaction that cleaves the polypeptide within the LPX_3X_4G motif of the sorting signal and covalently cross-links the amino-terminal portion of the cleaved polypeptide to the peptidoglycan to display the polypeptide on the surface of the Gram-positive bacterium.
- 69. A method for displaying a polypeptide on the surface of a Grampositive bacterium comprising the steps of:
- (a) expressing a polypeptide having a sorting signal at its carboxy—terminal end, the sorting signal having: (1) a motif of LPX₃X₄G therein; (2) a substantially hydrophobic domain of at least 31 amino acids carboxyl to the motif; and (3) a charged tail region with at least two positively charged residues carboxyl to the substantially hydrophobic domain, at least one of the two positively charged residues being arginine, the two positively charged residues being alocated at residues 31-33 from the motif, wherein X₃ is any of the twenty naturally-occurring L-amino acids and X₄ is selected from the group consisting of alanine, serine, and threonine;
 - (b) forming a reaction mixture including: (i) the expressed polypeptide; (ii) the substantially purified sortase—transamidase enzyme of claim 34; and (iii) a Gram—positive bacterium having a peptidoglycan to which the sortase—transamidase can link the polypeptide; and
 - (c) allowing the sortase-transamidase to catalyze a reaction that cleaves the polypeptide within the LPX₃X₄G motif of the sorting signal and covalently cross-links the amino-terminal portion of the cleaved polypeptide to the peptidoglycan to display the polypeptide on the surface of the Gram-positive bacterium.
 - 70. A method for displaying a polypeptide on the surface of a Grampositive bacterium comprising the steps of:
- (a) expressing a polypeptide having a sorting signal at its carboxy—
 terminal end, the sorting signal having: (1) a motif of LPX₃X₄G therein; (2) a substantially hydrophobic domain of at least 31 amino acids carboxyl to the motif; and
 (3) a charged tail region with at least two positively charged residues carboxyl to the substantially hydrophobic domain, at least one of the two positively charged residues being arginine, the two positively charged residues being located at residues 31-33
 from the motif, wherein X₃ is any of the twenty naturally-occurring L-amino acids and X₄ is selected from the group consisting of alanine, serine, and threonine:

- (b) forming a reaction mixture including: (i) the expressed polypeptide; (ii) the substantially purified sortase—transamidase enzyme of claim 35; and (iii) a Gram—positive bacterium having a peptidoglycan to which the sortase—transamidase can link the polypeptide; and
- (c) allowing the sortase-transamidase to catalyze a reaction that cleaves the polypeptide within the LPX₃X₄G motif of the sorting signal and covalently cross-links the amino-terminal portion of the cleaved polypeptide to the peptidoglycan to display the polypeptide on the surface of the Gram-positive bacterium.

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- 71. A method for displaying a polypeptide on the surface of a Grampositive bacterium comprising the steps of:
- (a) cloning a nucleic acid segment encoding a chimeric protein into a Gram-positive bacterium to generate a cloned chimeric protein including therein a carboxyl-terminal sorting signal, the chimeric protein including the polypeptide to be displayed, the sorting signal having: (1) a motif of LPX₃X₄G therein; (2) a substantially hydrophobic domain of at least 31 amino acids carboxyl to the motif; and (3) a charged tail region with at least two positively charged residues carboxyl to the substantially hydrophobic domain, at least one of the two positively charged residues being arginine, the two positively charged residues being located at residues 31-33 from the motif, wherein X₃ is any of the twenty naturally-occurring L-amino acids and X₄ is selected from the group consisting of alanine, serine, and threonine;
- (b) growing the bacterium into which the nucleic acid segment has been cloned to express the cloned chimeric protein to generate a chimeric protein including therein a carboxyl-terminal sorting signal; and
- (c) binding the polypeptide covalently to the cell wall by the enzymatic action of a sortase—transamidase expressed by the Gram—positive bacterium involving cleavage of the chimeric protein within the LPX₃X₄G motif so that the polypeptide is displayed on the surface of the Gram—positive bacterium in such a way that the polypeptide is accessible to a ligand.
- 72. A polypeptide displayed on the surface of a Gram-positive bacterium by covalent linkage of an amino-acid sequence of LPX_3X_4 derived from cleavage of an LPX_3X_4G motif, wherein X_3 is any of the twenty naturally-occurring L-amino acids and X_4 is selected from the group consisting of alanine, serine, and threonine, the polypeptide being displayed on the surface of the Gram-positive bacterium in such a way that the polypeptide is accessible to a ligand.

- 73. A covalent complex comprising:
- (a) the polypeptide of claim 72; and
- (b) an antigen or hapten covalently cross-linked to the polypeptide.

- 74. The covalent complex of claim 73 wherein an antigen is covalently cross-linked to the polypeptide.
- 75. The covalent complex of claim 73 wherein a hapten is covalently cross-linked to the peptide.
 - 76. A method for vaccination of an animal comprising the step of immunizing the animal with the displayed polypeptide of claim 72 to generate an immune response against the displayed polypeptide.

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77. A method for vaccination of an animal comprising the step of immunizing the animal with the covalent complex of claim 73 to generate an immune response against the antigen or hapten of the covalent complex.

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- 78. A method for screening for expression of a cloned polypeptide comprising the steps of:
- (a) expressing a cloned polypeptide as a chimeric protein having a sorting signal at its carboxy-terminal end, the sorting signal having: (1) a motif of LPX₃X₄G therein; (2) a substantially hydrophobic domain of at least 31 amino acids carboxyl to the motif; and (3) a charged tail region with at least two positively charged residues carboxyl to the substantially hydrophobic domain, at least one of the two positively charged residues being arginine, the two positively charged residues being located at residues 31–33 from the motif, wherein X₃ is any of the twenty naturally-occurring L-amino acids and X₄ is selected from the group consisting of alanine, serine, and threonine;

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(b) forming a reaction mixture including: (i) the expressed chimeric protein; the substantially purified sortase—transamidase enzyme of claim 1; and (iii) a Gram—positive bacterium having a peptidoglycan to which the sortase—transamidase can link the polypeptide through the sorting signal;

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(c) binding the chimeric protein covalently to the cell wall by the enzymatic action of a sortase-transamidase expressed by the Gram-positive bacterium involving cleavage of the chimeric protein within the LPX₃X₄G motif so

that the polypeptide is displayed on the surface of the Gram-positive bacterium in such a way that the polypeptide is accessible to a ligand; and

(d) reacting the displayed polypeptide with a labeled specific binding partner to screen the chimeric protein for reactivity with the labeled specific binding partner.

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- 79. A method for screening for expression of a cloned polypeptide comprising the steps of:
- (a) expressing a cloned polypeptide as a chimeric protein having a sorting signal at its carboxy-terminal end, the sorting signal having: (1) a motif of LPX₃X₄G therein; (2) a substantially hydrophobic domain of at least 31 amino acids carboxyl to the motif; and (3) a charged tail region with at least two positively charged residues carboxyl to the substantially hydrophobic domain, at least one of the two positively charged residues being arginine, the two positively charged residues being located at residues 31-33 from the motif, wherein X₃ is any of the twenty naturally-occurring L-amino acids and X₄ is selected from the group consisting of alanine, serine, and threonine;
- (b) forming a reaction mixture including: (i) the expressed chimeric protein; (ii) the substantially purified sortase—transamidase enzyme of claim 3; and (iii) a Gram—positive bacterium having a peptidoglycan to which the sortase—transamidase can link the polypeptide through the sorting signal;
- (c) binding the chimeric protein covalently to the cell wall by the enzymatic action of a sortase-transamidase expressed by the Gram-positive bacterium involving cleavage of the chimeric protein within the LPX₃X₄G motif so that the polypeptide is displayed on the surface of the Gram-positive bacterium in such a way that the polypeptide is accessible to a ligand; and
- (d) reacting the displayed polypeptide with a labeled specific binding partner to screen the chimeric protein for reactivity with the labeled specific binding partner.

80. A method for screening for expression of a cloned polypeptide comprising the steps of:

(a) expressing a cloned polypeptide as a chimeric protein having a sorting signal at its carboxy-terminal end, the sorting signal having: (1) a motif of LPX₃X₄G therein; (2) a substantially hydrophobic domain of at least 31 amino acids carboxyl to the motif; and (3) a charged tail region with at least two positively charged residues carboxyl to the substantially hydrophobic domain, at least one of the two

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positively charged residues being arginine, the two positively charged residues being located at residues 31-33 from the motif, wherein X_3 is any of the twenty naturally-occurring L-amino acids and X_4 is selected from the group consisting of alanine, serine, and threonine;

- (b) forming a reaction mixture including: (i) the expressed chimeric protein; (ii) the substantially purified sortase—transamidase enzyme of claim 31; and (iii) a Gram—positive bacterium having a peptidoglycan to which the sortase—transamidase can link the polypeptide through the sorting signal;
- (c) binding the chimeric protein covalently to the cell wall by the enzymatic action of a sortase-transamidase expressed by the Gram-positive bacterium involving cleavage of the chimeric protein within the LPX₃X₄G motif so that the polypeptide is displayed on the surface of the Gram-positive bacterium in such a way that the polypeptide is accessible to a ligand; and
- (d) reacting the displayed polypeptide with a labeled specific binding partner to screen the chimeric protein for reactivity with the labeled specific binding partner.
 - 81. A method for screening for expression of a cloned polypeptide comprising the steps of:
 - (a) expressing a cloned polypeptide as a chimeric protein having a sorting signal at its carboxy-terminal end, the sorting signal having: (1) a motif of LPX₃X₄G therein; (2) a substantially hydrophobic domain of at least 31 amino acids carboxyl to the motif; and (3) a charged tail region with at least two positively charged residues carboxyl to the substantially hydrophobic domain, at least one of the two positively charged residues being arginine, the two positively charged residues being located at residues 31-33 from the motif, wherein X₃ is any of the twenty naturally-occurring L-amino acids and X₄ is selected from the group consisting of alanine, serine, and threonine;
- (b) forming a reaction mixture including: (i) the expressed chimeric protein; (ii) the substantially purified sortase—transamidase enzyme of claim 32; and (iii) a Gram—positive bacterium having a peptidoglycan to which the sortase—transamidase can link the polypeptide through the sorting signal;
- (c) binding the chimeric protein covalently to the cell wall by the enzymatic action of a sortase-transamidase expressed by the Gram-positive bacterium involving cleavage of the chimeric protein within the LPX₃X₄G motif so that the polypeptide is displayed on the surface of the Gram-positive bacterium in such a way that the polypeptide is accessible to a ligand; and

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- (d) reacting the displayed polypeptide with a labeled specific binding partner to screen the chimeric protein for reactivity with the labeled specific binding partner.
- 82. A method for screening for expression of a cloned polypeptide comprising the steps of:
- (a) expressing a cloned polypeptide as a chimeric protein having a sorting signal at its carboxy-terminal end, the sorting signal having: (1) a motif of LPX₃X₄G therein; (2) a substantially hydrophobic domain of at least 31 amino acids carboxyl to the motif; and (3) a charged tail region with at least two positively charged residues carboxyl to the substantially hydrophobic domain, at least one of the two positively charged residues being arginine, the two positively charged residues being located at residues 31-33 from the motif, wherein X₃ is any of the twenty naturally-occurring L-amino acids and X₄ is selected from the group consisting of alanine, serine, and threonine;
- (b) forming a reaction mixture including: (i) the expressed chimeric protein; (ii) the substantially purified sortase—transamidase enzyme of claim 33; and (iii) a Gram—positive bacterium having a peptidoglycan to which the sortase—transamidase can link the polypeptide through the sorting signal;
- (c) binding the chimeric protein covalently to the cell wall by the enzymatic action of a sortase-transamidase expressed by the Gram-positive bacterium involving cleavage of the chimeric protein within the LPX₃X₄G motif so that the polypeptide is displayed on the surface of the Gram-positive bacterium in such a way that the polypeptide is accessible to a ligand; and
- (d) reacting the displayed polypeptide with a labeled specific binding partner to screen the chimeric protein for reactivity with the labeled specific binding partner.
- 83. A method for screening for expression of a cloned polypeptide comprising the steps of:
 - (a) expressing a cloned polypeptide as a chimeric protein having a sorting signal at its carboxy-terminal end, the sorting signal having: (1) a motif of LPX₃X₄G therein; (2) a substantially hydrophobic domain of at least 31 amino acids carboxyl to the motif; and (3) a charged tail region with at least two positively charged residues carboxyl to the substantially hydrophobic domain, at least one of the two positively charged residues being arginine, the two positively charged residues being located at residues 31-33 from the motif, wherein X₃ is any of the twenty naturally-

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occurring L-amino acids and X₄ is selected from the group consisting of alanine, serine, and threonine;

- (b) forming a reaction mixture including: (i) the expressed chimeric protein; (ii) the substantially purified sortase—transamidase enzyme of claim 34; and (iii) a Gram—positive bacterium having a peptidoglycan to which the sortase—transamidase can link the polypeptide through the sorting signal;
- (c) binding the chimeric protein covalently to the cell wall by the enzymatic action of a sortase-transamidase expressed by the Gram-positive bacterium involving cleavage of the chimeric protein within the LPX₃X₄G motif so that the polypeptide is displayed on the surface of the Gram-positive bacterium in such a way that the polypeptide is accessible to a ligand; and
- (d) reacting the displayed polypeptide with a labeled specific binding partner to screen the chimeric protein for reactivity with the labeled specific binding partner.

84. A method for screening for expression of a cloned polypeptide comprising the steps of:

- (a) expressing a cloned polypeptide as a chimeric protein having a sorting signal at its carboxy-terminal end, the sorting signal having: (1) a motif of LPX₃X₄G therein; (2) a substantially hydrophobic domain of at least 31 amino acids carboxyl to the motif; and (3) a charged tail region with at least two positively charged residues carboxyl to the substantially hydrophobic domain, at least one of the two positively charged residues being arginine, the two positively charged residues being located at residues 31-33 from the motif, wherein X₃ is any of the twenty naturally-occurring L-amino acids and X₄ is selected from the group consisting of alanine, serine, and threonine;
- (b) forming a reaction mixture including: (i) the expressed chimeric protein; (ii) the substantially purified sortase—transamidase enzyme of claim 35; and (iii) a Gram—positive bacterium having a peptidoglycan to which the sortase—transamidase can link the polypeptide through the sorting signal;
- (c) binding the chimeric protein covalently to the cell wall by the enzymatic action of a sortase-transamidase expressed by the Gram-positive bacterium involving cleavage of the chimeric protein within the LPX₃X₄G motif so that the polypeptide is displayed on the surface of the Gram-positive bacterium in such a way that the polypeptide is accessible to a ligand; and

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- (d) reacting the displayed polypeptide with a labeled specific binding partner to screen the chimeric protein for reactivity with the labeled specific binding partner.
- 85. A method for screening for expression of a cloned polypeptide comprising the steps of:
- (a) cloning a nucleic acid segment encoding a chimeric protein into a Gram-positive bacterium to generate a cloned chimeric protein including therein a carboxyl-terminal sorting signal, the chimeric protein including the polypeptide whose expression is to be screened, the sorting signal having: (1) a motif of LPX₃X₄G therein; (2) a substantially hydrophobic domain of at least 31 amino acids carboxyl to the motif; and (3) a charged tail region with at least two positively charged residues carboxyl to the substantially hydrophobic domain, at least one of the two positively charged residues being arginine, the two positively charged residues being located at residues 31-33 from the motif, wherein X₃ is any of the twenty naturally-occurring L-amino acids and X₄ is selected from the group consisting of alanine, serine, and threonine;
- (b) growing the bacterium into which the nucleic acid segment has been cloned to express the cloned chimeric protein to generate a chimeric protein including therein a carboxyl-terminal sorting signal;
- (c) binding the polypeptide covalently to the cell wall by the enzymatic action of a sortase—transamidase expressed by the Gram—positive bacterium involving cleavage of the chimeric protein within the LPX₃X₄G motif so that the polypeptide is displayed on the surface of the Gram—positive bacterium in such a way that the polypeptide is accessible to a ligand; and
- (d) reacting the displayed polypeptide with a labeled specific binding partner to screen the chimeric protein for reactivity with the labeled specific binding partner.
- 86. A method for the diagnosis or treatment of a bacterial infection caused by a Gram-positive bacterium comprising the steps of:
- (a) conjugating an antibiotic or a detection reagent to a protein including therein a carboxyl-terminal sorting signal to produce a conjugate, the carboxyl-terminal sorting signal having: (1) a motif of LPX₃X₄G therein; (2) a substantially hydrophobic domain of at least 31 amino acids carboxyl to the motif; and (3) a charged tail region with at least two positively charged residues carboxyl to the substantially hydrophobic domain, at least one of the two positively charged residues

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being arginine, the two positively charged residues being located at residues 31-33 from the motif, wherein X_3 is any of the twenty naturally-occurring L-amino acids and X_4 is selected from the group consisting of alanine, serine, and threonine; and

- (b) introducing the conjugate to an organism infected with a Grampositive bacterium in order to cause the conjugate to be sorted and covalently crosslinked to the cell walls of the bacterium in order to treat or diagnose the infection.
- 87. The method of claim 86 wherein an antibiotic is conjugated to the protein.
- 88. The method of claim 87 wherein the antibiotic is selected from the group consisting of a penicillin, ampicillin, vancomycin, gentamicin, streptomycin, a cephalosporin, amikacin, kanamycin, neomycin, paromomycin, tobramycin, ciprofloxacin, clindamycin, rifampin, chloramphenicol, norfloxacin, and a derivative of these antibiotics.
- 89. The method of claim 86 wherein a detection reagent is conjugated to the protein.
- 90. A conjugate comprising an antibiotic or a detection reagent covalently conjugated to a protein including therein a carboxyl-terminal sorting signal to produce a conjugate, the carboxyl-terminal sorting signal having: (1) a motif of LPX₃X₄G therein; (2) a substantially hydrophobic domain of at least 31 amino acids carboxyl to the motif; and (3) a charged tail region with at least two positively charged residues carboxyl to the substantially hydrophobic domain, at least one of the two positively charged residues being arginine, the two positively charged residues being located at residues 31-33 from the motif, wherein X₃ is any of the twenty naturally-occurring L-amino acids and X₄ is selected from the group consisting of alanine, serine, and threonine.
 - 91. The conjugate of claim 90 wherein an antibiotic is conjugated to the protein.
- 92. The conjugate of claim 91 wherein the antibiotic is selected from the group consisting of a penicillin, ampicillin, vancomycin, gentamicin, streptomycin, a cephalosporin, amikacin, kanamycin, neomycin, paromomycin,

tobramycin, ciprofloxacin, clindamycin, rifampin, chloramphenicol, norfloxacin, and a derivative of these antibiotics.

- 93. The conjugate of claim 90 wherein a detection reagent is conjugated to the protein.
 - 94. A composition comprising:
 - (a) the conjugate of claim 90; and
 - (b) a pharmaceutically acceptable carrier.

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- 95. A substantially purified protein having at least about 50% match with best alignment, in at least one subunit of the protein, with the amino acid sequences of at least one of the putative *Bacillus* peptidase (SEQ ID NO: 3), the aminopeptidase P of *Lactococcus lactis* (SEQ ID NO: 4), or the proline dipeptidase of *Lactobacillus delbrueckii lactis* (SEQ ID NO: 5) and having sortase—transamidase activity.
- 96. The substantially purified protein of claim 95 wherein the match with best alignment with the amino acid sequences of at least one of the putative *Bacillus* peptidase (SEQ ID NO: 3), the aminopeptidase P of *Lactococcus lactis* (SEQ ID NO: 4), or the proline dipeptidase of *Lactobacillus delbrueckii lactis* (SEQ ID NO: 5) is at least about 60%.
- 97. The substantially purified protein of claim 96 wherein the match with best alignment with the amino acid sequences of at least one of the putative Bacillus peptidase (SEQ ID NO: 3), the aminopeptidase P of Lactococcus lactis (SEQ ID NO: 4), or the proline dipeptidase of Lactobacillus delbrueckii lactis (SEQ ID NO: 5) is at least about 70%.
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- 98. A substantially purified protein having sortase—transamidase activity and a hydrophobicity profile of at least one subunit of the protein, that, determined as the mean absolute value of the hydrophobicity difference per residue, differs from the hydrophobicity profile of a putative *Bacillus* peptidase (SEQ ID NO: 3) by no more than about 2 units on the hydrophobicity scale.

- 99. The substantially purified protein of claim 98 wherein the hydrophobicity profile differs from the hydrophobicity profile of the putative *Bacillus* peptidase (SEQ ID NO: 3) by no more than about 1 unit.
- 5 100. The substantially purified protein of claim 99 wherein the hydrophobicity profile differs from the hydrophobicity profile of the putative *Bacillus* peptidase (SEQ ID NO: 3) by no more than about 0.5 unit.
- 101. A nucleic acid sequence encoding the substantially purified protein of claim 95.
 - 102. A nucleic acid sequence encoding the substantially purified protein of claim 98.
- 103. A vector comprising the nucleic acid sequence of claim 101 operatively linked to at least one control sequence that controls the expression or regulation of the nucleic acid sequence.
- 104. A vector comprising the nucleic acid sequence of claim 102 operatively linked to at least one control sequence that controls the expression or regulation of the nucleic acid sequence.
 - 105. A host cell transfected with the vector of claim 103.
- 25 106. A host cell transfected with the vector of claim 104.

- 107. A method for producing a substantially purified protein having sortase-transamidase activity comprising the steps of:
- (a) culturing the host cell of claim 105 under conditions in which the host cell expresses the protein having sortase-transamidase activity; and
- (b) purifying the expressed protein to produce substantially purified protein having sortase-transamidase activity.
- 108. A method for producing a substantially purified protein having sortase—transamidase activity comprising the steps of:
 - (a) culturing the host cell of claim 106 under conditions in which the host cell expresses the protein having sortase—transamidase activity; and

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- (b) purifying the expressed protein to produce substantially purified protein having sortase-transamidase activity.
- 109. A substantially purified protein having sortase—transamidase

 5 activity and a hydrophobicity profile of at least one subunit of the protein, that,
 determined as the mean absolute value of the hydrophobicity difference per residue,
 differs from the hydrophobicity profile of the sequence of SEQ ID NO: 31 by no more
 than about 2 units on the hydrophobicity scale.
- 110. The substantially purified protein of claim 109 wherein the hydrophobicity profile differs from the hydrophobicity profile of the sequence of SEQ ID NO: 31 by no more than about 1 unit.
- 111. The substantially purified protein of claim 110 wherein the
 hydrophobicity profile differs from the hydrophobicity profile of the sequence of SEQ
 ID NO: 31 by no more than about 0.5 unit.
 - 112. A nucleic acid sequence encoding the substantially purified protein of claim 109.

113. A vector comprising the nucleic acid sequence of claim 112 operatively linked to at least one control sequence that controls the expression or regulation of the nucleic acid sequence.

114. A host cell transfected with the vector of claim 113.

- 115. A method for producing a substantially purified protein having sortase-transamidase activity comprising the steps of:
- (a) culturing the host cell of claim 114 under conditions in which the host cell expresses the protein having sortase-transamidase activity; and
 - (b) purifying the expressed protein to produce substantially purified protein having sortase-transamidase activity.

RIEDDIL-ITENGCOVETKCTKDLIVLT*

344

RIEDD-LVITKTGCQVLTLAPKELIVL-*

(SEQ ID No: 2 &

_	Match	ing	Matching Percentage (Total Window: 21%, Alignment Window:	22%)
_	S.aureus	9-	DPKYQ	43
	S.pyogenes	Н	MLQYSQKLPKEFAMSGFLEQRLGHCLRQMAEKGLEALLVTHLTNSYYLTG	20
		44	IVLESQMKAINEIR-P	93
		51	FSGTAATVLITAKRVLITDSRYTLLAKASVEGFDIIESRTPLKVVAELL	100
		94		143
	•	101	EADQIDCLGFEDQVSFSFYQAMQAELSGITLLAQSGFVEHLRLIKDASEI	150
		144	DAISSK	193
		151		200
		194	GYGKE	243
		201	IIVASGYLSAMPHGRASDKVIQNKESLTMDFGCYYNHYVSDMTRTIHIGQ	250
	- •	244	IG-LEIHEGPM-LAR-T	293
		251		300
	- •	294	IQDKL-QVNNCVIVEPGVYIEGLGGI	343
		301	HGIGHGIGLDIHENPFFGKSEQLLQAGMVVTDEPGIYLDNKYGV	350

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Mat	ching	Matching Percentage (Total Window: 46%, Alignment Window: 47%)	(Total	Window:	468,	Alignment	Window:	478)
S.aureus	ا	-3DPKLKEIYQIVLESQMKA-INEIRPGMTGAEADAISRNYLESKG 46	YOIVLES	OMKA-INEI	RPGMT(SAEADAISRN)	TLESKG	46
S.pyogenes	249	249 GQVTDEER-EIYALVLAAN-KALIAKASAGMTYSDFDGIPRQ-LITEA-G 298	YALVLAA	N-KALIAKA:	SAGMT	rsdedeipro-	LITEA-G	298
	47	47 YGKEFGHSLGHGIGLEIHEGPMLARTIODKLOVNNCVTVEPGVYIEG	SIGLEIH	EGPMLARTI(PDKLQ-	VNNCVTVE	SPGVYIEG	96
	299	299 YGSRFTHGIGHGIGLDIHENPFFGKSEQL-LQAGMVVTDEPGIY 348	SIGLDIH	I I Enpffgkse(707-70	AGMVVTDE	PGIY	348
	97	97 LGGIRIEDDIL-ITENGCQVFTKCTKDLIVLT*	DIL-ITE	NGCOVETKC:	TKDLIV	/LT*	•	146
	349	349 LDNKYGVRIEDD-LVITKTGCQVLTLAPKELIVL-*		TGCQVLTLA	PKELIV	72-*	•	398

(SEQ ID No: 2 & 34)

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)	
•	;		
	_		

600 ATCATTTGAT	590 CCAGCGGCAC	550 550 560 570 580 590 600 CTGGCTAATT TTTTAGATT TCGTATGCGT CAGTATGGTG CCAGCGGCAC ATCATTTGAT	570 TCGTATGCGT	560 TTTAGATTT	550 CTGGCTAATT	
540 TGAACGTGAC	520 530 540 ATTAAACCAG GGACAACCAC TGAACGTGAC	520 ATTAAACCAG	510 TCTTGATTTT	490 500 510 SACAAAGCAT TTGAAGATGC TCTTGATTTT	490 GACAAAGCAT	
480 CTCGATCTCA	470 CTAAAGCGTG	430 440 450 460 460 470 480 TTACGTCTTA TTAAGGACGC CTCTGAAATC GATACCATTG CTAAAGCGTG CTCGATCTCA	450 CTCTGAAATC	440 TTAAGGACGC	430 TTACGTCTTA	
420 Tgtggagcat	410 AGTCAGGTTT	370 380 390 400 410 420 GCCATGCAAG CAGAACTGTC AGGAATAACC TTGCTTGCTC AGTCAGGTTT TGTGGAGCAT	390 AGGAATAACC	380 CAGAACTGTC	370 GCCATGCAAG	
360 TTTTTACCAG	340 350 360 360 360 360 360	340 GAGGACCAGG	330 CCTTGGTTTT	310 320 GAGGCTGATC AAATAGATTG	310 GAGGCTGATC	
300 Agaattgtta	290 AGGTTGTGGC	270 280 290 300 cgaaagccgc accccctta accttctta	270 CGAAAGCCGC	250 260 GTTGAGGGAT TTGATATTAT	250 GTTGAGGGAT	
240 CCTTGCTTGC TAAAGCTAGT		220 TCACGTTATA	210 GATCACAGAT	190 200 ACGGCCAAAC GTCGTGTTT	190 ACGGCCAAAC	
170 CTGCAGCAAC TGTTTTGATA		160 TTTTCTGGAA	150 CTTGACAGGT	130 CATTTAACCA ATAGTTATTA	130 CATTTAACCA	
120 TCTAGTCACC	110 TAGAGGCTCT	70 80 90 100 110 CGATTAGGTC ACTGCCTAAG GCAGATGGCA GAGAAGGGGC TAGAGGCTCT	90 GCAGATGGCA	80 ACTGCCTAAG	70 CGATTAGGTC	
60 TTTAGAACAA	50 TGTCAGGATT	10 20 30 40 50 60 AIGCTACAAT ATTCTCAAAA GTTACCAAAG GAGTTCGCGA TGTCAGGATT TTTAGAACAA	30 GTTACCAAAG	20 ATTCTCAAAA	10 ATGCTACAAT	

660	720	780	840
TGACAAGGTT	CTATGTTAGT	GATTTATGCT	GACTTATAGT
650	710	770	830
GACGCGCCAG	ACTACAATCA	AAGAACGTGA	GCGCTGGCAT
640	700	760	820
ATGCCTCATG	TTTGGGTGTT	GTTACTGATG A	GCTAAAGCTA
630	690	740	810
TCTCTCTGCC	GACCATGGAC	CATTCA TATTGGCCAA GTTA	GGCTTTAATT
610 620 630 640 650 660 ATCATTGTAG CTTCAGGCTA TCTCTGCC ATGCCTCATG GACGCGCCAG TGACAAGGTT	670 680 690 700 710 720 ATCCAGAATA AAGAGAGCTT GACCATGGAC TTTGGGTGTT ACTACAATCA CTATGTTAGT	SGACC	790 830 810 820 830 840 CTTGTTCTTG CTGCTAATAA GGCTTTAATT GCTAAAGCTA GCGCTGGCAT GACTTATAGT
610	670	730	790
ATCATTGTAG	ATCCAGAATA	GATATGACGA	CTTGTTCTTG

900 TCGCTTCACA	960 TGGGAAATCT	1020 TTTGGATAAC	1080 TCAAGTCTTG	1140
850 860 870 880 890 900 GACTTTGACG GTATGGCAG TCGCTTCACA	910 920 930 940 950 CATGGCATTG GTCATGGCAT CGGGCTTGAC ATCCATGAGA ATCCATTTT	970 980 1000 1010 1020 GAGCAACTIC ICCAAGCIGG AAIGGIGGIA ACAGAIGAGC CAGGIAICTA ITIGGAIAAC	1030 1040 1050 1060 1070 1080 AAATATGGTG TCCGTATTGA AGATGACTTG GTTATCACAA AAACTGGTTG TCAAGTCTTG	1130
880 ACTGAGGCGG	940 ATCCATGAGA	1000 ACAGATGAGC	1060 GTTATCACAA	1120 TAA
870 CCAACTCATC	930 CGGGCTTGAC	990 Aatggtggta	1050 Agatgacttg	1090 1100 1110 ACCTTGGCAC CCAAAGAATT AATTGTATTG TAA.
860 GTATTCCGCG	920 GTCATGGCAT	980 TCCAAGCTGG	1040 TCCGTATTGA	1100 CCAAAGAATT
850 GACTTTGACG	910 CATGGCATTG	970 GAGCAACTTC	1030 AAATATGGTG	1090 ACCTTGGCAC

(SEQ ID No: 33 & 34)

5900 TTA	5954 GCT 	6008 ACT	6062 CGT	6116 CGC 	6170 CTT
			TCA S		
			GAT		
5891 TCA	5945 GGG 	5999 TTT 	6053 ACA 	6107 ATC 	6161 ATA
ATG	AAG	66T	ATC	ATT	S O
800 800	GAG	ACA T	TTG	GAT 	GAT
5882 TTC 		n,	6044 GTT		
GAG 	ATG	TAC	CGT R	66A	GAG
AAG K			CG1		_
5873 CCA 	5927 AGG			6089 GTT 	
			225 ₹		
			ACG	•	_
5864 CAA 	5918 CAC 	5972 TTA		6080 AAA	6134 GTG
TCT			TTG	GCT	
TAT 	TTA	ACC	GTT	CIT	AAG K
5855 CAA 	5909 CGA	5963 GTC 	6017 ACT 	6071 TTG	6125 CTT
			GCA		
ATG	GAA	CIT	GCA	TAT	ACG

FIG. 4

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6224 CTG	6278 ATT	6332 AAA 	6386 GAC 	TCA TCA	6494 GCC 	34)
GAA 	CTT	GAC	CGT	ACA T	00 00 1 %	Š
GCA A	CGT	TCA	GAA 	9	GGA	(SEQ. ID
6215 CAA	6269 TTA 	6323 ATC 	6377 ACT 	6431 AGC 	6485 CAT	S)
ATG	CAT	TCG	ACC	GCC	CCT	
80C	GAG 	TGC	ACA	GGT	ATG	
6206 CAG	6260 GTG 	6314 GCG 	6368 GGG	6422 TAT	6476 GCC 	
TAC	TIT	AAA K	CCA	CAG	TCT	
TTT	GGT 	GCT	AAA	CGT	CIC	
6197 TCT 	6251 TCA 	6305 ATT 	6359 ATT 	6413 ATG 	6467 TAT 	
TIT	CAG	ACC	TTT 	CGT) 1 - 1 - 9	
TCG	GCT	GAT 	GAT D	TTT 	TCA	
6188 GTA 	6242 CTT 	6296 ATC 	6350 CTT 	6404 GAT	6458 GCT 	
CAG	TTG	GAA	GCT	TTA	GTA 	
GAC	ACC	TCT	GAT	TTT	ATT	
6179 GAG 	6233 ATA 	6287 GCC 	6341 GAA 	6395 AAT 	6449 ATC 	
				GCT		
6GT	TCA	AAG	GCA P	CTG	TT.	

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6548	TAC	*	6602 ACT	H	6656 ATT	Н	6710 CAA	a	6764 GGC
	TGT	ပ	GTT	>	TTA	1 -1	၁၅၁	æ	CAT
	999	v	CAA	0	GCT	4	922	۵,	661 0
5539	TTT	[24	5593 GGC	9	3647 AAG	×	3701 ATT	H	6755 ATT
•	GAC	۵	ATT	H	AAT	z	GGT	ဗ)))))
	ATG								CAT
	ACC								6746 ACA T
	TTG		ACC	6	CTT	1 -1	GAC	Ω	TTC
	AGC		AGG						CGC 8
			6575 ACG	=	5629 CTT	1	5683 TAT	>	6737 AGT
	AA:	×	ATG	Σ	GCT	A	ACT	H	၂ ၁
	AAT	z	GAT	۵	TAT	,	ATG	Σ	TAT
	CAG	ø	5566 AGT	w	6620 ATT	Н	5674 GGC	ប	6728 GGT
	ATC	H	GTT	>	GAG	ы	GCT	æ	929
									GAG
									6719 ACT
	GAC	۵	AAT	z	GAA	ம	AAA	×	ATC
	AGT	တ	TAC	 	GAT	۵	GCT	Æ	CTC

FIG. 4

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6818		1	0 r r		6872	AAA	AA	6872 AAC AAA TAT N K Y	AAA	AAA K	AAA K	AAA X X X X X X X X X X X X X X X X X X	AAA K	AAA K
809	GAG		ы		GAT A	-	Ω	917						
9	TCT	1	N P F F G K S E	Ψ	TTG	1	L D	v	GGT TGT	1	ပ			
	AAA	1	×		TAT		V T D E P G I Y L		ACT	1	←			
6800	999		ပ	6854	ATC	1	H	8069	AA	1	×		3,	
	TIT		(E4		GGT	i	ဗ		ACA		₽		TAA	
	TTT	!	Ĺų		CCA	!	Δ,		ATC		H		TTG	
6791	დ	1	۵,	6845	GAG	1	ы	6689	GTT	1	>	6953	GTA	
	AAT	1	z		GAT	1	Ω		TTG	1	н		ATT	
	GAG	1	ы		ACA		H		GAC	1	۵		TTA	
6782	C ATC CAT GAG	1	Ħ	6836	GTA		T > >	6890				6944	GAA	
	ATC	1	H		GTG	1	>		GAA	1	ല		AAA	
	SA	į	۵		ATG	1	Σ		ATT	1	н		ပ္ပ	
6773	CII		I G L	6827	GGA	1	ပ	6881	CGT	1	æ	6935	ACC TIG GCA CCC AAA GAA TIA AIT GIA IIG IAA	
	999		ဗ		CAA GCT	1	4		GIC		> গ		TIG	
	ATC	1	H		CAA	1	a		GGT		ဖ		ACC	

	dfryteqakv	daelvpvaes	mrsqgadsss		mevhespgls	111	
	isgdkaafit	tyasysavis	iavanelefy	csditrtvav	fghstghglg		(SEQ ID No: 3)
	qftgsaglav	gfeqnsmtyg	ltfmkpgise	ldfgayykgy	hiaakgygdy	ddivitengn	(SEQ
	tsntnvØmt	tvesfgikrl	kladdafrhi	kliesgdlvt		ipetggvrie	
	gqlgidgmli	ggsligttad	eeikileeaa	sslphgvasd	gvahikpgmt	mvvtvepgiy	
	mkleklrnlf	qvkgfeiieh	veklrlikss	fdmivasglr	yqvvfdaqal	vrssailepg	
	-	61	121	181	241	301	
210110							1

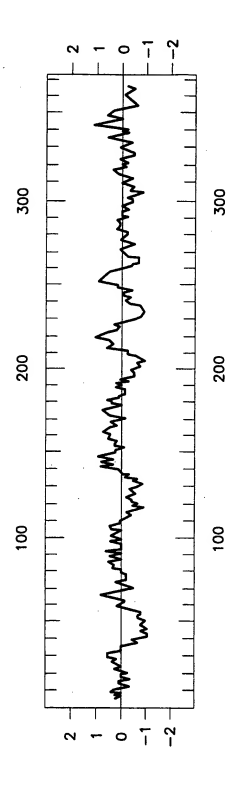


FIG. 6

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dleasgisfe vddkmrtiye

vanfldfkmr

vheipyfngs

dmtrtifvgs hgighglgld vltkapkeli

fgcyyehyas ekadfggyft dllvtengce

qydniprevi pefggvried

vitdepgiyl

mtenglrsgm

dsrysemarg

ltqkrnifmt krlskaatkl

gfsgtagtvf feetvdyaff flepgrtele

tdmknifylt sasesvknma adeafmsalr iqfgdpvtid

dlfstsnfvl

ORIGIN

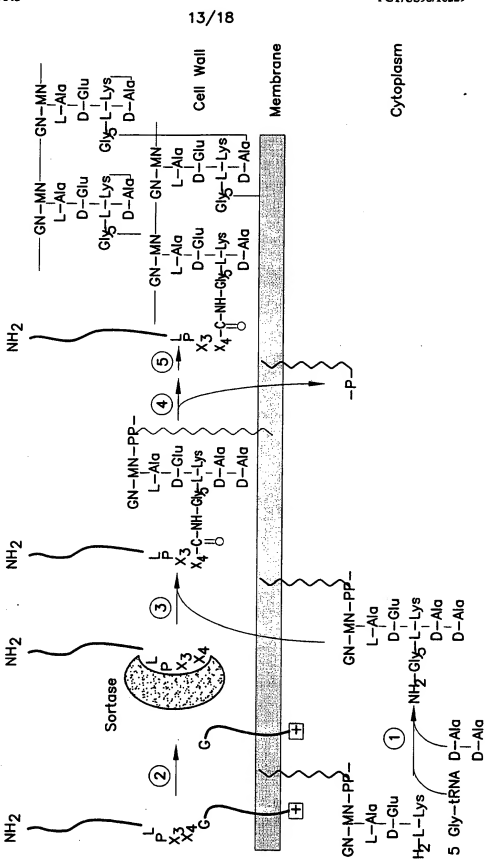
lphgvatskm kqvkagmtya ltenids111 rdpislltel islikkacei mrieklkvkm elrqikdese tivasgkrss liknfeiiet tvrkaneali 61 121 181 241 301

ID No: 4) (SEQ

ORIGIN

ravvsqieyq yasdssrtva ýfihrlghgi fkpfthtske pflfcpalny ygalhaqfpd kiitdagyge edcgvltkdg fklfafkdae vekngltvah fealrngvte lfdlgtmheg 3 (SEQ. ID No: <u>qfit</u>dpeeri krtkdygnwa eeadfafqig mntvqpnelv taseldgvar yipgfagvri sspttinyft pwskiaeeik naanphqgps aaidaakpgm selvklrkag gmcfsiepgi qengmdvayv dvvgyldsed iahirlfkte sfdtivqagk iyevnrtaqq angndvvlee eeakasawdg sdfskdlsdf gmevhefpsi lkvlpvke mnldklgnwl ygeptdkmre lklqkgvmqt 1 61 121 181 241 301 361

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60	120	180	240	300	360	420	480	540
ATTAATCGTA	GAAAATGTTN	AGCCGTATAT	GCTGACGAAA	ATTTTAACTG	CAAATGCTAN	Agaggtgcat	TACATTANAT	TTGCNATTGG
10 20 30 40 50 60 $-ff$ -CGACTC TAGAGGACAA GCAACTAAGC AGGCGCCAAA TTATGAAATT ATTAATCGTA	70 80 90 100 AATCTACTAT TATTGGTGAG ATTAAAAAT TGCTACNCCA ANAAAATTTT	150 160 170 180 NTTTATGATA CATACCTTGA ATTAAATAAA AGCCGTATAT	190 200 210 230 CATTAATAAG CATTTCTAAT ACTGTAGATA AAATTAGAGA CGTCCAAGAT	250 260 270 280 290 300 TIGCTTTAAT TCAAAAAGCA GCTAATATTG TTGATGAAAC ATATGAATAT ATTTTAACTG	310 320 330 340 350 360 TIGTAAAAGC NNGCATGACT GAAAAANAAT TAAAGGCAAT ATTAGAAAGC CAAATGCTAN	420 CGATTGINGC ATCTGGTCCT AGAGGTGCAT	460 ATTGAAAAG GCGACATGAT	490 530 510 520 530 540 TITGGGCGCN TINITATAAC CGGCTATIGI ICAAATITIA CIANAACAII ITGCNAIIGG
40	100	160	220	280	340		460	520
AGGCGCCAAA	TGCTACNCCA	CATACCTTGA	AAATTAGAGA	TTGATGAAAC	Taaaggcaat		Attgaaaag	TCAAATTTTA
30	90		210	270	330	390	450	510
GCAACTAAGC	ATTAAAAAAT		ACTGTAGATA	GCTAATATTG	Gaaaaanaat	TCTTTCGATA	TGATAAAATT	CGGCTATTGT
20	80	130	200	260	320	370 380	430 440	500
Tagaggacaa	Tattggtgag	GTTTTGAGG GCATCATGTN	CATTTCTAAT	TCAAAAAGCA	NNGCATGACT	AATTAGGAGC AGATGGACCN	TACCACCATG GTGTTGCAAG	TINTTATAAC
10	70	130	190	250	310	370	430	490
-/-cgactc	AATCTACTAT	GTTTTGAGGG	CATTAATAAG	TIGCTTTAAT	TTGTAAAAGC	AATTAGGAGC	TACCACCATG	TTTGGGCGCN

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(SEQ ID No: 28 & 29)	OEO ID				
	•		•		TTTAA
1080	1070	1060	1050	1040	1030
1020	1010	1000	990	980	970
Agaagagata	GTATAAGAAT	GGTTTGGGCG	TNATATAGAA	NACCTGGTGT	GTTACAGTAG
960	950	940	930	920	910
TAACAACTGT	AACTTCAAGT	ATACAAGATA	GGNTCGTACG	GGCCAATGCT	ATCCATGAAG
900	890	880	870	860	850
TGGTTTAGAA	GACATGGTAT	CATTCACTAG	AGAATTTGGA	GGTATGGAAA	GAGTCAAAAG
840	830	820	810	800	790
AAACTATTTA	CCATTTCAAG	GAAGCTGATG	GACTGGTGCA	GACCTGGCAT	AATGAGATTA
780	770	760	750	730 740	730
Gaaagcaatt	AATCTCAAAT	ATAGTACTTG	AATATATCAA	CCAGATCCTA AACTGAAAGA	CCAGATCCTA
720	710	700	690	680	670
Tattggagaa	GAACATTIGC	GATATTACTA	CTATTGTTCA	ATTATAACGG	TINGGCGCGT
660	650	640	630	620	610
NACATTAGAT	GCGACATGAT	ATTGAAAAAG	TGATAAAATT	GTGTTGCAAG	TTACCACATG
600	590	570	570	560	550
Tagaggtgca	CATCTGTTCA	TGAAAGAA- <i>ff</i> -ATTGTAG	TGAAAGAA-	ATCCNTAAAC	GAAAAACCCG

State CCT Anh CTG Anh CTG Anh CTG CT							•	
CCT	54 GCA 	108 AGA	162 CAT 	216 GAT 	270 GGT 	324 CAA O		
GAT CCT AAA CTG AAA GAA ATA TAT CAA ATA CAA CAA ATA CAA CAA ATA CAA CAA CAA ATA CAA CAA <th>AA ×</th> <td>TCA</td> <td>868 6</td> <td>SA O</td> <td>GAA</td> <td>TGT</td> <td></td> <td>~</td>	AA ×	TCA	868 6	SA O	GAA	TGT		~
GAT CCT AAA CTG AAA GAA TAT CAA ATA CTA ATA CTA ATA CTA ATA CTA GAA ATA CTA CAA ATA CTA CAA ATA CTA CAA ATA CTA CTA <th>ATG</th> <td>ATT</td> <td>CTA</td> <td>ATA </td> <td>ATA </td> <td>GGT</td> <td></td> <td></td>	ATG	ATT	CTA	ATA 	ATA 	GGT		
GAT CCT AAA CTG AAA CAA ATA TAT CAA ATA CAA ATA CAA ATA CAA ATA CAA ATA CAA ATA CAT CAA CAT CAT <th>CAA CAA</th> <td>99 GCC </td> <td>153 TCA </td> <td>207 ACG</td> <td>261 TAT </td> <td>315 AAT </td> <td>ě</td> <td><u></u></td>	CAA CAA	99 GCC 	153 TCA 	207 ACG	261 TAT 	315 AAT 	ě	<u></u>
GAT CCT AAA CTG AAA GAA ATA TAT CAA ATA TAT CAA ATA ATA ATA ATA ATA GTA ATA ATA ATA CTA ATA ATA CTA ATA ATA <th>TCT</th> <td>GAT</td> <td>CAT</td> <td>CGT R</td> <td>GTT </td> <td>GAA</td> <td>TAA </td> <td>(SEC</td>	TCT	GAT	CAT	CGT R	GTT 	GAA	TAA 	(SEC
GAT CCT AAA CTG AAA GAA ATA TAT CAA ATA GAA ATA CAA ATA CAA ATA CAA ATA CAA ATA CAA ATA ATA CAA ATA CAA ATA CAA ATA CAA ATA ATA CAA ATA ATA <th>GAA </th> <td>GCT P A</td> <td>868 0</td> <td>GCT</td> <td>66T 6</td> <td>ACA T</td> <td>ACA</td> <td></td>	GAA 	GCT P A	868 0	GCT	66T 6	ACA T	ACA	
GAT CCT AAA CTG AAA GAA ATA TAT CAA ATA T P K L K E I Y Q I ATT GAG ATT AGA CCT GGC ATG ACT GGC ATG GGT ACT GGC ATG ACT GGC AAA GGC CCA AAA GGC AAA GGC AAA GGC CCA AAA GGC CCA AAA AAA AAA GGC AAA AAA AAA AAA AAA AAA AAA AAA	36 CTT 	90 GAA	144 TTT 	198 CTG	252 CCT 	306 ATT 	360 TTA 	
GAT CCT AAA CTG AAA GAA ATA TAT CAA I P K L K E I Y Q ATT GAG ATT AGA CCT GGC ATG ACT ATT GAG ATT AGA CCT GGC ATG ACT AAA TTA GAG TTA AGA TTA GGA ATG AGA ATG AGA AGG ATG AGG AGA AGG AGG AGA AGG	GTA	GCA A	GAA 	ATG	GAA	TTA 	GTT	
GAT CCT AAA CTG AAA GAA ATA ATA TAT I P K L K E I Y I AAT GAG ATT AGA CCT GGC ATG AAC TAA GAG ATT AGA CCT GGC ATG AAC TAA GAG TTA GAG TAA GGC TAT GGT TTA GAG TTA GAA ATC GAA ATC AAA TTA GAG TTA GAA ATC GAA ATC GGT T T T T T T T TTG T T T T T T T T GGT T T T T T T T T T T T T T T T T T T<	ATA 	GGT	AAA X	CCA	GTA 	ATT	ATA	
GAT CCT AAA CTG AAA GAA ATA D P K L K E I ATT AAT GAG ATT AGA CCT GGC I N E I R F G G AAC TAT GAG TCA AAAA GGC GGC GG GG AAC TAT GAG TCA AAA GGG GG I B I B I GG GG I I B I I GG GG I I I I I I I I I I I I I I I I I	CAA	ACT T	135 GGA 	189 GGG 	243 ACA T	297 GAT	351 CTT	
GAT CCT AAA CTG AAA GAA GAA L L K L K E E C C C C C C C C C C C C C C C C C	TAT	ATG	TAT 	GAA E	GTT	GAT	GAC	
GAT CCT AAA CTG AAA C D P K L K L K 63 ATT AGA G I N E I R I N E I R I N E I R GGT ATT GGT TTA GAA GG I G L E S AAA CTT CAA GTT AGA K L Q V N TTG GGC GGT ATA AGA A L G G I R I R I R I R I G I R I R I G I R I R I G I R I R I G I R I R I G I R I R I G I R I G I R I R I G I R I G I R I R I G I R I R I G I R I R I G I R I G I R I R I G I R I R I G I R I R I G I R I R I G I R I R I G G I R I R I G G I R I R I G G I R I R I G G I R I R I G G I R I R I G G I R I R I G G I R I R I G G I R I R I G G I R I R I G G G G G I R I R I G G G G G G G I R I R I G G G G G G G G G G G G G G G G G G G	ATA I	9	999	CAT H	TGT 	GAA	AAA	
GAT CCT AAA CTG D P K L 1	GAA E	CCT CCT	126 AAA 	180 ATC 	234 AAC 	288 ATA 	342 ACA 	
GAT CCT AAA GAG I I I I I I I I I I I I I I I I	AAA X	AGA 	TCA	GAA	AAC	AGA R	7GC	
GAT CCT D D P D P T R T R T R T R T R T R T R T R T R T	CTG 	ATT	GAG	TTA 	GTT 	ATA 	AAA X	
GGT AAA AAA V	AAA K	63 GAG	117 TTA 	171 GGT 	225 CAA 	279 GGT G	333 ACT 	
	CCT	AAT	TAT	ATT 	CTT	၁၅၅	TTT 	
în	GAT	ATT	AAC	66T 6	AAA	TTG	GTC 	
	ດ໌							

FIG.

17/18

	•					
3311 GCG Ala	3365 ATC Ile	3419 AAA Lys	3473 GAT	3527 ATT Ile	3581 CAA Gln	31)
ATA Ile	GGC G1y	CGT Arg	ATA Ile	AAA Lys	AAG Lys	30 & 31)
GAA G1u	GAA Glu	AGT Ser	TAT Tyr	AAC 	CCT	ë
3302 GTA Val	3356 GTC Val	3410 GTT 	3464 ATA Ile	3518 GCA Ala	3572 GAA Glu	3' (SEQ ID
AAA Lys	GAA Glu	AAA Lys	66C 61y	ACT	ATA Ile	3' (SE(
GGT	TCG	GAA Glu	GAT Asp	AAA Lys	GCG Ala	TAA
3293 TTA Leu	3347 ACT Thr	3401 TTA Leu	3455 GAA Glu	3509 TCA	3563 ACA Thr	3617 AAG Lys
AAA Lys	GCT	AAT	AAA Lys	ATT 11e	ATG	GAA Glu
TCA	ATA Ile	ACA Thr	AGT Ser	AAT Asn	AAT Asn	ATT Ile
3284 AAT Asn	3338 AGT Ser	3392 GAA Glu	3446 GAG Glu	3500 GTT 	3554 TCT Ser	3608 GTT Val
TCA	GCA	AAA Lys	ATC Ile	GGT Gly	ATT Ile	ATC 11e
TAT	ATT Ile	TTA Leu	AAA Lys	CAT	TCA	CAA Gln
3275 GAT ASP	3329 GTT 	3383 GAA Glu	3437 TTA 	3491 AAA 	3545 AAT Asn	3599 ACA Thr
ACT	TCT	GCT Ala	GAT Asp	TTA	TTT Phe	ATT
GTA Val	CTA	TTT	CGT Arg	GCA 	ATT Ile	CAC His
3266 AAA Lys	3320 GTG 	3374 CAT	3428 AGC Ser	3482 TGT	3536 TCA Ser	3590 ATT Ile
GTC	GAA Glu	GGC G1y	TTA	TAT	ACG	AAT
ATG Met	CCA 	ACT Thr	AAT Asn	GTA 	CAA	ATT Ile
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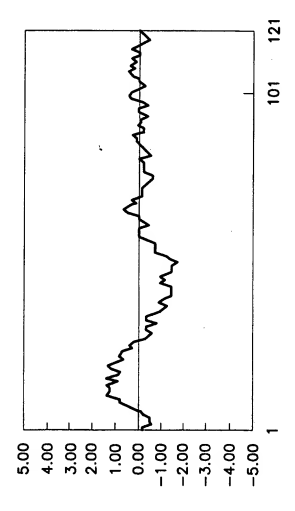


FIG. 1.

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                                     25
                                                         30
      Ala Ile Ser Arg Asn Tyr Leu Glu Ser Lys Gly Tyr Gly Lys Glu Phe
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                                 40
                                                      45
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                              55
                                                  60
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130 135 140
10
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145 150 155 160
       Ile Ala Val Ala Asn Glu Leu Glu Phe Tyr Met Arg Ser Gln Gly Ala
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                                           170
                                                                 175
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180 185 190
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210 215 220
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225 230 235 240
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       Tyr Gln Val Val Phe Asp Ala Gln Ala Leu Gly Val Ala His Ile Lys
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      Phe Glu Ile Ile Glu Thr Arg Asp Pro Ile Ser Leu Leu Thr Glu Leu
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                           70
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      Ser Ala Ser Glu Ser Val Lys Asn Met Ala Phe Glu Glu Thr Val Asp
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                                            90
      Tyr Ala Phe Phe Lys Arg Leu Ser Lys Ala Ala Thr Lys Leu Asp Leu 100 105 110
       Phe Ser Thr Ser Asn Phe Val Leu Glu Leu Arg Gln Ile Lys Asp Glu
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               115
                                   120
      Ser Glu Ile Ser Leu Ile Lys Lys Ala Cys Glu Ile Ala Asp Glu Ala
130 135 140
      Phe Met Ser Ala Leu Arg Phe Ile Glu Pro Gly Arg Thr Glu Ile Glu
                          150
                                              155
      Val Ala Asn Phe Leu Asp Phe Lys Met Arg Asp Leu Glu Ala Ser Gly
165 170 175
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185

His Gly Val Ala Thr Ser Lys Met Ile Gln Phe Gly Asp Pro Val Thr

190

180

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                                                   235
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       Ala Asp Phe Gly Gln Tyr Phe Thr His Gly Ile Gly His Gly Leu Gly 275 280 285
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       Gln Leu Arg Ser Gly Met Val Ile Thr Asp Glu Pro Gly Ile Tyr Leu
305 310 315 320
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                           310
                                                 315
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                                          25
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       Gln Asn Trp Ala Val Glu Lys Asn Gly Leu Thr Val Ala His Tyr Gln
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       Ala Leu His Ala Gln Phe Pro Asp Ser Asp Phe Ser Lys Asp Leu Ser
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130 135 140
       Lys Leu Arg Lys Ala Gly Glu Glu Ala Asp Phe Ala Phe Gln Ile Gly 145 150 155 160
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       Phe Glu Ala Leu Arg Asn Gly Val Thr Glu Arg Ala Val Val Ser Gln
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                                           170
       Ile Glu Tyr Gln Leu Lys Leu Gln Lys Gly Val Met Gln Thr Ser Phe
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       Thr Ala Gln Gln Ala Ala Ile Asp Ala Ala Lys Pro Gly Met Thr Ala
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                                         265
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       Ser Glu Leu Asp Gly Val Ala Arg Lys Ile Ile Thr Asp Ala Gly Tyr
275 280 285
       Gly Glu Tyr Phe Ile His Arg Leu Gly His Gly Ile Gly Met Glu Val
290 295 300
       His Glu Phe Pro Ser Ile Ala Asn Gly Asn Asp Val Val Leu Glu Glu
65
                           310
                                                 315
       Gly Met Cys Phe Ser Ile Glu Pro Gly Ile Tyr Ile Pro Gly Phe Ala
325 330 335
                                              330
                                                                     335
       Gly Val Arg Ile Glu Asp Cys Gly Val Leu Thr Lys Asp Gly Phe Lys
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Ala Ile Asn Glu Ile Arg Pro Gly Met Thr Gly Ala Glu Ala Asp Ala
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		ata Ile	tt 1	taa													1017
15																	
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	Ala	Ile	Ser 35	Arg	Asn	Tyr	Leu	Glu 40		Lys	Gly	Tyr	Gly		Glu	Phe	
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		Val	Xaa	Pro	Gly 85		Xaa	Ile	Glu	Gly 90		Gly	Gly	Ile	Arg 95		
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			100>														
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				gaa Glu													96
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J J	gtc Val	gaa Glu	ggc Gly 35	atc Ile	act Thr	ggc Gly	cat His	ttt Phe 40	gct Ala	gaa Glu	tta Leu	aaa Lys	gaa Glu 45	aca Thr	aat Asn	tta Leu	144
60				agt													192
	Glu	Lys 50	Val	Ser	Arg	Lys	Asn 55	Leu	Ser	Arg	Asp	60	ьуs	lle	Glu	Ser	
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				aaa Lys													288

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      Val Glu Gly Ile Thr Gly His Phe Ala Glu Leu Lys Glu Thr Asn Leu
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                                                    45
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      Glu Lys Val Ser Arg Lys Asn Leu Ser Arg Asp Leu Lys Ile Glu Ser
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                             55
                                                 60
      Lys Glu Asp Gly Ile Tyr Ile Asp Val Tyr Cys Ala Leu Lys His Gly
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                                              75
      Asn Ile Ser Lys Thr Ala Asn Lys Ile Gin Thr Ser Ile Phe Asn Ser
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                                                              95
                     85
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      Phe Leu Glu Gln Arg Leu Gly His Cys Leu Arg Gln Met Ala Glu Lys
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      ggg cta gag gct ctt cta gtc acc cat tta acc aat agt tat tac ttg
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      Gly Leu Glu Ala Leu Leu Val Thr His Leu Thr Asn Ser Tyr Tyr Leu
                                   40
               35
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J	cgt Arg 65	gtt Val	ttg Leu	atc Ile	aca Thr	gat Asp 70	tca Ser	cgt Arg	tat Tyr	acc Thr	ttg Leu 75	ctt Leu	gct Ala	aaa Lys	gct Ala	agt Ser 80	240
10	gtt Val	gag Glu	gga Gly	ttt Phe	gat Asp 85	att Ile	atc Ile	gaa Glu	agc Ser	cgc Arg 90	acg Thr	ccg Pro	ctt Leu	aag Lys	gtt Val 95	gtg Val	288
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		gac Asp															480
30		aaa Lys															528
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45		gcc Ala 210															672
,,,		agc Ser															720
50		atg Met							His		Gly						768
55		cgt Arg															816
60		aaa Lys															864
65		caa Gln 290															912
65		ggt Gly															960

-11-

	aaa to Lys Se	gag c Glu G	aa ctt In Leu 325	ctc Leu	caa Gln	gct Ala	gga Gly	atg Met 330	gtg Val	gt .	aaca	gatg	ag		1005
5	ccaggta aaactg	tct at ctt gt	ttggat caagtc	aa ca tt ga	aata	atgg:	t gte	ccgta	attg gaat	aag taa	atga ttgt	ctt d	ggtt. gtaa	atcaca	1065 1119
10	•	210> 3 211> 3 212> P 213> S	32	cocci	ıs P	yogei	nes								
15	Met Leu 1		yr Ser 5					10					15	•	
	Phe Let	2	0				25					30		-	
20	Gly Let Thr Gly	35				40					45		_		
20	50 Arg Val				55					60			-	-	
	65			70					75			_		80	
25	Val Glu	GIÀ 5	ne Asp 85	116	ше	GIU	Ser	Arg 90	Thr	Pro	Leu	Lys	Val 95	Val	
	Ala Glu		eu Glu 00	Ala	Asp	Gln	Ile 105	Asp	Суз	Leu	Gly	Phe 110	Glu	Asp	
	Gln Val	Ser P. 115	he Ser	Phe	Tyr	Gln 120	Ala	Met	Gln	Ala	Glu 125	Leu	Ser	Gly	
30	Ile Thr 130		eu Ala	Gln	Ser 135	Gly	Phe	Val	Glu	His 140	Leu	Arg	Leu	Ile	
	Lys Asp 145			150	-				155		_			160	
35	Asp Lys		165					170				_	175		
	Thr Glu	1	80				185					190		_	
	Gly Ala	195	_			200					205	_	-		
40	Ser Ala 210		ro His	Gly	Arg 215	Ala	Ser	Asp	Lys	Val 220	Ile	Gln	Asn	Lys	
	Glu Ser 225	Leu T	hr Met	Asp 230	Phe	Gly	Cys	Tyr	Tyr 235	Asn	His	Tyr	Val	Ser 240	
45	Asp Met	Thr A	rg Thr 245	Ile	His	Ile	His	Ile 250		Gln	Val	Thr	Asp 255		
	Glu Arg			Ala	Leu	Val	Leu 265		Ala	Asn	Lys	Ala 270		Ile	
	Ala Lys	_	• -	Gly	Met	Thr 280		Ser	Asp	Phe	Asp 285		Ile	Pro	
50	Arg Gln 290	Leu I	le Thr	Glu	Ala 295		Tyr	Gly	Ser	Arg 300		Thr	His	Gly	
	Ile Gly		ly Ile	Gly 310		Asp	Ile	His	Glu 315		Pro	Phe	Phe	Gly 320	
55	Lys Ser	Glu G	ln Leu 325		Gln	Ala	Gly	Met 330	_	Val					

INTERNATIONAL SEARCH REPORT

International application No. PCT/US98/16229

A. CLASSIFI IPC(6) :C12N	CATION OF SUBJECT MATTER		
US CL :435/1	7 9/10, 15/63, 1/21, 1/15, 1/19; C12Q 1/48; (93, 320.1, 252.3, 254.11, 15, 7.32; 536/23.2	C07H 21/04	•
According to Inter	mational Patent Classification (IPC) or to bot	h national classification and IPC	
	BARCHED		
Minimum docume	ntation searched (classification system follow	red by classification symbols)	
	73, 320.1, 252.3, 254.11, 15, 7.32; 536/23.2		
Documentation sea	arched other than minimum documentation to t	he extent that such documents are included	in the fields searched
Electronic data bas Please See Extra	se consulted during the international search (s Sheet.	name of data base and, where practicable	, search terms used)
C. DOCUMEN	ITS CONSIDERED TO BE RELEVANT	· · · · · · · · · · · · · · · · · · ·	
Category* C	itation of document, with indication, where a	ppropriate, of the relevant passages	Relevant to claim No.
g/:4	abase PROMT on STN, Inform 120001, 'Antibacterials, SIGA Wement.' abstract. R & D Focus I de abstract.	yeth Ayerst, SIGA licensing	1-49, 57-71 and 95-115
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Further docu	ments are listed in the continuation of Box C	See patent family annex.	
Special categ	cories of cited documents:	"T" later document published after the inte	mational filing date or priority
document de to be of part	fining the general state of the art which is not considered inular relevance	date and not in conflict with the appli the principle or theory underlying the	
earlier docum	nent published on or after the international filing date	"X" document of particular relevance; the	claimed invention cannot be
CHARGE ED MINE	high may throw doubts on priority claim(s) or which is blish the publication date of another citation or other in (as specified)	considered novel or cannot be consider when the document is taken alone "Y" document of particular relevance: the	·
•	ferring to an oral disclosure, use, exhibition or other	considered to involve an inventive combined with one or more other such	step when the document is documents, such combination
document put the priority d	blished prior to the international filing date but later than	being obvious to a person skilled in the "&" document member of the same patent	
ate of the actual c	ompletion of the international search	Date of mailing of the international sea	rch report
02 NOVEMBER	1998	19 NOV 1998	•
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Box PCT Washington, D.C. 2		KAWALLAU FOR	
	3) 305-3230	Telephone No. (703) 308-0196	
	second sheet)(July 1992)*	(103) 308-0170	

INTERNATIONAL SEARCH REPORT

International application No. PCT/US98/16229

Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1. Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
2. Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This International Searching Authority found multiple inventions in this international application, as follows:
1. As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:
4. X No required additional search fees were timely paid by the applicant. Consequently, this international search report is
restricted to the invention first mentioned in the claims; it is covered by claims Nos.: 1-49, 57-71 and 95-115
Remark on Protest The additional search fees were accompanied by the applicant's protest.
No protest accompanied the payment of additional search fees.

INTERNATIONAL SEARCH REPORT

International application No. PCT/US98/16229

B. FIELDS SEARCHED		
F75		

Electronic data bases consulted (Name of data base and where practicable terms used):

APS, STN: PROMT, CAPLUS, CEN, MEDLINE, BIOSIS, BIOTECHDS, CANCERLIT, CEABA, DRUGNL, EMBASE, FSTA, SCISEARCH, TOXLINE, PIRA, WPIDS, EUROPATFULL search terms: sortase, sort, enzyme, transamidase, gram, positive

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